

pplicants: J.G. BEDNORZ ET AL. : Date: March 29, 1988

Filed: 05/22/87 : Serial No.: 06/053,307

Group Art Unit: 115 : Examiner: Dennis Albrecht

FOR: NEW SUPERCONDUCTIVE COMPOUNDS HAVING HIGH TRANSITION TEMPERATURE, AND METHODS FOR THEIR USE AND PREPARATION

DECLARATION OF ALBERT M. TORRESSEN WITH RESPECT TO HIGH TO SUPERCONDUCTIVITY

Commissioner of Patents and Trademarks Washington, D.C. 20231

Sir:

- I, Albert M. Torressen, hereby declare and say that:
- 1. I joined the International Business Machines Corporation (IBM Corp.) on June 12, 1978, and began my employment as a laboratory technician at the Thomas J. Watson Research Center, Yorktown, New York. In 1986, I was a senior lab specialist at the Watson Research center, and reported to Dr. Stephan von Molnar. My specialty has been measurement of thermal and transport properties of a material including, for example, measurements of heat capacity, resisitivity, and Hall coefficients. Generally, these are dynamic measurements which serve to charactize a material.
- 2. On approxmately October 22, 1986, Richard L. Greene approached Stephan von Molnar to enlist his assistance in obtaining specific heat measurements of samples which he said were new superconducting materials that had been recieved from Georg Bednorz and Alex Mueller of IBM's Zurich research laboratory. These were ceramic oxide materials comprised of Ba-La-Cu-O. In turn, von Molnar asked me to assist Greene in making these measurements, since I had expertise in the use of the apparatus and had done similar measurements for many years. I began to calibrate the apparatus and to prepare for specific heat

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measurements at that time, and worked continuously on a daily basis, from about October 22, 1986 through the remainder of 1986 and into 1987 on these measurements. Richard L. Greene was in the laboratory during these measurements and supervised the experimentation, relying on my expertise with respect to the apparatus. Later, we analyzed the data together.

- 3. Richard Greene wanted to do specific heat measurements of these superconducting samples. Such measurements are used to dertermine if a material is a superconductor and also to tell how much of the material is superconducting. In general, the specific heat of the apparatus is first carefully measured using a bolometer in order to provide background specific heat and to calibrate the apparatus. After this, the actual sample to be measured is attached to the bolometer and the specific heat of the entire apparatus, including the sample, is again measured. When the background specific heat (due to the bolometer) is subtracted, the specific heat of the sample can be determined. This is done over a temperature range, in our measurements 2-50K, in order to obtain a plot of specific heat versus temperature for the sample being measured. This is a dynamic measurement in which we look at how the heat in the sample decays as a function of time through a known heat leak and from that extrapolate via a computer program specific heat versus temperature. This is a known procedure that is done in many laboratories.
- 4. The commercial bolometer that Greene and I used to measure the specific heat of these superconducting Zurich samples was comprised of an insulating aluminum oxide on which strips of RuO₂ were evaporated. Electrical contacts were made to the RuO₂ strips, each strip having two AuCu wires attached thereto which were in turn connected to an ambient temperature control (about 2K). A heat pulse was then applied to the bolometer and its temperature decay versus time was measured. This was done over the aforementioned temperature range in order to get background specific heat and to calibrate the apparatus. Generally, it takes us about two days to measure the background specific heat in order to prepare for the actual specific heat measurements of the sample.

 Measurement of each sample also takes about two days, so approximately four days represents the total time required for each measurement. This is based on full time activity.

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- 5. Prior to making the specific heat measurements, the samples are prepared. The superconducting oxide samples received from Bednorz and Mueller were pellet samples which were then cut into slices by a diamond wheel. I believe that this was done by Richard Greene. To make the specific heat measurements, a sample having a mass of about 20 mg is required, it being desirable to have a flat surface sufficient to provide good thermal contact. In the calculation of specific heat, the weight (mass) of the sample is factored into the equation. Also, I made specific checks of the measuring apparatus throughout this time period (October 22, 1986 February, 1987) and I made continual checks on the measuring apparatus in order to ensure its calibration. All of the data collected during the measurements was also provided to a personal computer that was interfaced to the apparatus. In this manner, a computer printout of all of the measurements was available.
- 6. Exhibit A attached hereto is comprised of copies of eight pages of my laboratory notebook and a copy of the notebook cover entitled "Specific heat - Zurich oxide - 10/21/86". All of the writing of these pages was entered by me on the dates indicated on the pages. These pages are true copies of the corresponding pages of my laboratory notebook that I have numbered in red in the upper right hand corner to enable discussion of the data on each page. As I mentioned previously, background specific heat and calibration of the instrumentation was done prior to mounting the sample and measuring the specific heat of the sample. For example, the data on the top two-thirds of page one shows the measured data for the calibration and background measurements. bottom of this page indicates the specific heat data that was obtained October 27, 1986 when the sample was mounted on the RuO, bolometer. weight of the sample is also mentioned, the sample being designated "BLCO 2 - I". This stands for Ba-La-Cu-O ceramic superconductor. I noted that it was a "multi-phase sample" which is not a particular good choice for a specific heat measurement. Generally, the data is more clean and easy to interpret if the sample contains only a single phase. Measurements on the apparatus and with this particular sample continued on October 28 and 29, 1986, the data that Greene and I obtained being listed on page 2 on Exhibit A.

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- 7. On November 3, 1986 we used a new sample, in this case the sample designated "BLCO 21 -II". We used a new calibration of the bolometer and then mounted the sample on the bolometer to take the data, which was obtained on November 4, 5, 6, 1986. The data for this sample measurement is contained on pages 4 and 5 of Exhibit A, where it is noted that this is a single phase sample. Additional data is also contained on page 6 of Exhibit A.
- 8. Richard Greene and I analyzed this data between about November 10 and November 19, 1986. At that time, the samples showed only a very small (1 2 percent) anomaly at the transition temperature, which was sufficiently small that no obvious bump occurred in this specific heat data. Because these materials had a very broad transition in resistivity versus temperature, such a very small effect was expected.
- 9. On or about November 19, 1986, the apparatus was changed slightly by installing a capacitor and mounting a magnet on the specific heat cryostat. The instrument was then recalibrated and measured, the data of this being shown on page 7 of Exhibit A. A capacitance bridge was used to control temperature as noted on page 8 of Exhibit A which showed further data taken on the instrumentation. After this time, we continued to take specific heat measurements of additional samples of the Ba-La-Cu-O oxide superconductors obtained from Bednorz and Mueller.
- 10. Exhibit B is a true copy of pages of a printout from the personal computer that was interfaced with the apparatus used to make the aforementioned specific heat measurements. These pages illustrate the background measurements and calibration of the instrumentation, as well as the data that were obtained when the sample was mounted on the bolometer. Additionally, many plots are included in this Exhibit which are plots of the data that were measured. Specific heat measurements of the samples are plotted where the samples are designated either "Zurich oxide" or "ZO". Sometimes the sample is also designated "Zurich oxide BLCO, etc." In December, 1986, additional samples were obtained from Bednorz and Mueller, these new samples containing Sr instead of Ba. They are designated "SLCO", which represents Sr-La-Cu-Oxide. The heat capacity of these samples was also measured, as represented by the plot dated January 9, 1987. Measurements were made in the absence of and in the presence of an applied magnetic field H. Sometimes the plots show

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the sample plus the designation "BG". This indicates that background was also present in the measurement.

- 11. The data that we obtained during our specific heat measurements are representative of the type of data which we now obtain on refined samples of these superconducting copper oxide materials. At the time we made our initial measurements in 1986, we were somewhat puzzled by the small vertical offset that occurred for temperatures extrapolated from 2-0K. However, such offsets have been found to be a characteristic of the superconducting copper oxides of the type first discovered by Bednorz and Mueller.
- 12. In addition to his specific heat measurements, Richard Greene also measured resistivity versus temperature in the presence of a magnetic field, for these Ba-La-Cu-O samples. This was done in my lab, and I explained my experimental instruments and set-up to Greene. I observed Richard Greene making these measurements and saw the shift in resisitivity versus temperature curve with an applied magnetic field. This shift clearly indicated the superconducting nature of these samples at temperatures in excess of 30°K. The measurements described in this paragraph occurred in the last week of November, 1986 and in the first week of December, 1986. These measurements and our specific heat measurements were part of our continuous daily effort, from about October 22, 1986 to establish the superconductive properties of these samples above 300K. I recall these measurements clearly and remember Richard Greene asking me how to better stabilize the sample temperature while the applied magnetic field was changed in amplitude.
- 13. In addition to the acts described hereinabove relating to work done by Richard Greene and by me, I was aware of the work being done by Chang C. Tsuei to measure resistivity versus temperature for these Ba-La-Cu-O superconducting samples. I was present in the laboratory with Chang C. Tsuei and and Sung Il Park during their measurement of at least one of these samples, and saw the hardcopy of a resistivity versus temperature plot developed by the xy recorder connected to their measurement apparatus. This plot showed the onset of superconductivity at approximately 35K followed by a broad transition to zero resistivity. I knew the nature of their experimentation and understood the data. This

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measurement and my observation occurred approximately the first week of November, 1986.

- 14. My recollection of the events described hereinabove is vivid, as there was great excitement about the importance of the discovery of new superconducting materials by Bednorz and Mueller. Because of this, activity continued on a daily basis, both morning and evening, to characterize these materials in coordination with Bednorz and Mueller.
- 15. All acts referred to hereinabove performed by myself, Richard L. Greene, Chang C. Tsuei, and Sung Il Park occurred in the United States.
- 16. I further declare that all statements made hereinabove are of my own knowledge and are true and that all statements made on information and belief are believed by me to be true. Further, I declare that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of a Patent Application or any patent issuing thereon.

Alleut m. Tonessen

ALBERT M. TORRESSEN

DATE: 31 March 1988

TORKESEN CHIBIT A

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Historia.

Do calibrations and Background Specific Heat

0216HTCB, DAT (5-33°K) } Ruo2 Celibrations 0216HTCB, DAT (24-40°K)

0226BGHD.DAT (6-40°K) Bochground Decay.

9 Bul + Wires + Grease

0226BGHE.DAT (6-10°K) 0226BGHG.DAT (6-25°K)

0226BGHH. DAT (10-20°K) 0226BGHI-DAT (10-40°K)

0226BGHJ. DAT (20-40K)

TXT

XT

TXT

10/23/86 0236PCH. DAT (To=6°K, Power Collection to 40°K) 0236LTC. DAT (Temperature Collection 2-12°K) 0236PCL. DAT (To=2°K, PC to 11°K) 0236BGL. DAT (To=2°K, 2-11°KBG)

5 Ample put IN - BLCO 2-I - Multiphase sample (bad choice Weight

0 276 PC H. DAT (To = 241t, Power Calib. to ~50t)

Ly Block 186.35R = 24th Carbo Glass Te

0 276 DAT (Decay 35-24th)

0776 ZOHB. DAT (Decay 40-24/K)

16/28/86

RUN Some More on DATA & Power Calibrations

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TXT 0286 PCL. DAT ( PC 2-8H)

TXT 0286 ZOLA. DAT ( Decay 2-8H)

TXT 0286 ZOLA. DAT ( Decay 12-5H)

TXT 0286 ZOHA. DAT ( PC 5-12H)

TXT 0286 ZOHB. DAT ( Decay 20-10H)

TXT 0286 ZOHC. DAT ( PC 10-20H)

TXT 0286 ZOHC. DAT ( Decay 28-17H)

TXT 0286 ZOHC. DAT ( PC 28-17H)
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- complex analyze date

he start some setting up

Charleton Absolute vide of the and only ages 200

10/29/86

TXT 0296HTC-DAT - Bolometer Calab at HT (28.50K)

029620 HA. DAT (Decay 30-24K)

029620 HB. DAT (Decay 33-24K)

029620 HC. DAT (Decay 37-24K) C.PAT Mechanis II mar a 43-24 decay

029620 HD. DAT (Decay 43-24K) 300 mm TXT file OK (37-24)

029620 HE. DAT (Decay 43-24K) -9 1 sec T

> Same base as Pum BGLB

11/3/86

New Sample + Background - hopefully single phase sample this
BLCO21-II time

Start with background + new bolometer calibration

Using 116 preamp on PAR - noise greatly reduced (see curves)

TXT \ N\$36 HTCA.DAT (5-32K) Ruoz calibration
TXT \ N\$36 HTCB_DAT (26-40K) "

TXT H \$36 BGHA. DAT (Decay 15-5 H)
TXT H \$36 PCHA. DAT (Power Calil: 5+15)

N\$36 BGHB.DAT (Deray 25 124)

11/4/86

Pump to ~2k and do LT runs

TXT NO46 LT CA-DAT (2-10H) Ruoz Calibration

C maybe 0 not 0 it is

TYT. NOY6 BGLA. DAT (Decay -8-18t) -> seemed to have long decay at end

TXT NO46BGLB. DAT (Dacay-8-2K) Rerumenth delle bourge T-still seem to see duft-bod smelle

TXT NO 46 PCLA. DAT (Power Calib. 18 - 84)

Back to 4.21 BAth

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THY NO 46 BGHA. DAT (Decay 25-121)

THY NO 46 PC HA. DAT (Power Calib 12-25K) -- > sook at this data

(NO 46 PC HB. DAT (never if HA) > better - the beginning

Use this one

TXT NO 46 BG HB. DAT (Decay 35-201)
TXT NO 46 PC HC. DAT (Pour Cult 20-351)

11/6/86

e e

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B

P

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Put on New Sample
                           BLCO 21-II
                                            Weight 14.82 mg
              (Single Phase)
                                                   ±.02mg
                                            2,5,12,20
                                                       BulV
                         ( Decay 15-5 t)
      NOGE ZOHA. DAT
                                                      9,1300
TXT ( NO 66 ZOHB . DAT
                         (Some 15-5k with T=100m)
     40 6 6 30 HC. DAT
                        ( Decay 25-12K ) T=180ms
      N$6670HD.DAT
                     ( Deary 35-20 K) 9=100mg
      1 TXT N P 66 HTCA. DAT (Bol. Calibration 5-100 K)
NO 66 HTCB. DAT
                        ( Bol. Calibration 26-40 H)
 TXT { N D 66 20 HE - DAT ( Decay 33-5K) T=103M)
1 TXT N $6670 HF. DAT ( Decay 12-6K) T= 100ms
1 TXT NO 6620 HE DAT ( same decay but put in front of cume)
1 TXT (NO 66 20 HA. DAT ( Decay 24-61)
1 TXT ( NO 6620 HI. DAT ( Decay 35-64)
1 TXT (NO66 PC HA DAT ( Power Calib 6-35 K)
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11/7/86

NOTELTCA DAT

(Tealit 2-10K)

N 076 ZOLA . DAT

(Neway 10-24)

NO 76 20 LB. DAT

(Same deeg - mu pts before end)

NOTEPELA. DAT (Power Cal. 2-10 K)

m7W510 SHEAT
Bordyourd from SEP24) BG9246
SVMC3, SVMT3 (2-12°K)

Barbyound from Nov4 CBG2K1, TBG2K1 (2-8°K)

Zurich OxIDE (BLC021 + BED) CAG, TAG (2-12°K)

Installed CS-4016R-B Copacitor 19 Nov. 1986

Mountail 5 Tesla magnet on Spenfix heat cryostate Cooled overreget to INa temperature

-77K 7.2611 V 120.72 6.341 NF. (SET ESCHOO @ 12.682 NF)

-- 4.2k 9.2015 1027.2 4.867 NF (set to will @ 1.000 nf)

Russel temperature to 6°K with LR-130 (Power) ~6°K 8.8762 600.1 (1.090 HF) 0 7.7308 " 1.140 54043 V

> NI9620HA. DAT (6-32°K) Zero Field 1.090 efter lecey

6°K 8.8765 600.1 (10.0918) 0 (using iR-130) 8.8780 600.2 10.0882 0 "CSC400

11-20-86

Wind Persistant heat switch

~6x 8.9830 601.5 (10.0882) 0 mm csc400

8 Costrolling @ ~ 6 k with Cycouttone Bridge (58400) (10.0880 NF) 11-21-86 RUOZ C2329 8.9224 606.1 (±0.3) 8.9214 605.9 5 MV (5A) 8.9204(t.001) 605.9(±1.0) 10 MV 8.9197 605.8 20 MV 55 K684° 8,9189 607,8 636.76/A 30MV 8.9178 604.8 YOMU 607.9 8.9152 50MV 8.9138 609.3 60 MV 70 A 636.7 G/A 8.9105 609.5 70MU 44.5690 KG. el-130 Cotrolle pour mogretouristance of CD329 monde

-6° 8.8977 600.0 70MU 0 -32° 7.76 " 540M 1.2804V

N21620HA. DAT (44 Tesus)

8.91 600. 0 0 7.76 " " 540m

N21620HB. DAT (Zero TCSU)

Celculate 70 MR & 2.00 Possible chong Ourte field.

TOPLESSEN |
WHIBIT B

1 76.57E-01 27.98E+00 28.00E+00 0.100 2 76.38E-01 28.92E+00 27.00E+00 0.100 3 76.19E-01 29.91E+00 30.00E+00 0.100 4 76.00E-01 30.90E+00 31.00E+00 0.100 5 75.82E-01 32.08E+00 32.00E+00 0.100 6 75.49E-01 34.03E+00 34.00E+00 0.100 75.17E-01 36.06E+00 36.00E+00 0.100 9 75.17E-01 37.06E+00 37.00E+00 0.100 10 74.87E-01 37.90E+00 38.00E+00 0.100 11 74.72E-01 38.93E+00 39.00E+00 0.100	Pt #	RUO2 VOLTS	CS35	SET TEMP	TEMP DEV	
	3 4 5 6 7 8 9 10	76.38E-01 76.19E-01 76.00E-01 75.82E-01 75.65E-01 75.49E-01 75.33E-01 75.17E-01 75.02E-01 74.87E-01	28.92E+00 29.91E+00 30.90E+00 32.08E+00 33.06E+00 34.03E+00 35.03E+00 36.06E+00 37.06E+00	29.00E+00 30.00E+00 31.00E+00 32.00E+00 33.00E+00 34.00E+00 35.00E+00 37.00E+00 38.00E+00	0.100 0.100 0.100 0.100 0.100 0.100 0.100 0.100	

'Ctrl/____' to end, 'Ctrl/F1' to locate, or 'Cursor Keys' to scroll.

Pt #	RUO2 VOLTS	C2327 TEMP	SET TEMP	TEMP DEV	
1	88.52E-01	49.80E-01	50.00E-01	0.050	·
3	86.92E-01 85.65E-01	59.78E-01 69.79E-01	60.00E-01	0.050 0.050	C ^B
4	84.62E-01	79.71E-01	80.00E-01	0.050	Ì
5	83.74E-01	89.67E-01	90.00E-01	0.050	
6	82.99E-01 82.33E-01	99.68E-01 10.98E+00	10.00E+00 11.00E+00	0.050 0.050	
á	81.72E-01	12.01E+00	12.00E+00	0.050	,
9	S1.19E-01	13.01E+00	13.00E+00	0.050	·
10	80.74E-01 80.29E-01	13.96E+00 14.99E+00	14.00E+00 15.00E+00	0.050	
12		15.98E+00	16.00E+00	0.050 0.050	
13		16.97E+00	17.00E+00	0.050	
14 15	1	17.97E+00	18.00E+00	0.050	
16		18.97E+00 19.97E+00	19.00E+00 20.00E+00	0.050 0.050	· .
17	78.23E-01	20.97E+00	21.00E+00	0.050	
18		22.03E+00	22.00E+00	0.050	
19 20	77.49E-01 77.45E-01	22.97E+00 23.99E+00	23.00E+00 24.00E+00	0.050 0.050	THE LETTER

3001 37 13								
"Utrl/i"	to	end.	Utr1/F1	to locate,	OF	Cursor	Kevs'	to scroll.

Pt #	RUO2 VOLTS	C2329 TEMP	SET TEMP	TEMP DEV	
21 22 23 24 25 26 27 28 29	77.21E-01 77.00E-01 76.77E-01 76.56E-01 76.36E-01 76.16E-01 75.99E-01 75.64E-01	24.97E+00 25.96E+00 26.97E+00 28.03E+00 29.00E+00 30.05E+00 31.00E+00 31.98E+00	25.00E+00 26.00E+00 27.00E+00 28.00E+00 29.00E+00 30.00E+00 32.00E+00 33.00E+00	0.050 0.050 0.050 0.050 0.050 0.050 0.050	

'Ctrl/____' to end, 'Ctrl/F1' to locate, or 'Cursor Keys' to scroll.

```
Min Temp to Plot
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                              K
 Max Temp to Plot
                     = 14.00
                              K.
 Min Volts to Plot = 8.00
                             Volts
 Max Volts to Plot = 10.0
                             Volts
 Plot Title
                     = RUO2 VS TEMP
 X Axis Plot Label = Sample Temp (K)
 Y Axis Plot Label = RU02 (Volts)
 Start Temperature = 002 K
 Temperature Step
                    = 0.5 \text{ K}
 Temperature Stop
                     = 014 K
 LR400 Range
                     = 20000 \text{ Dhms}
 Temperature Dev.
                    = 0.050 \text{ K}
 Wait Time
                 = 0010 seconds
 Notebook messages = 1 4
 RUO2 Calibration vs C2329 Carbon Glass Resistor
 4 WIRE CONFIGURATION FOR BOLOMETER
BACKGROUND MEASUREMENT
 STEPPING UP
 Number of points =
                              Values per point =
                      21
   RUO2 VOLTS
                    C2329 TEMP
                                        SET TEMP
                                                          TEMP DEV
    99.26E-01
                      19.96E-01
                                        20.00E-01
                                                            0.050
    96.15E-01
                     25.17E-01
                                        25.00E-01
                                                            0.050
    94.16E-01
                     29.68E-01
                                        30.00E-01
                                                            0.050
    92.41E-01
                     34.72E-01
                                        35.00E-01
                                                            0.050
    91.03E-01
                     39.68E-01
                                        40.00E-01
                                                            0.050
    89.85E-01
                     44.67E-01
                                        45.00E-01
                                                            0.050
    88.85E-01
                     49.68E-01
                                        50.00E-01
                                                            0.050
    87.98E-01
                     54.76E-01
                                        55.00E-01
                                                            0.050
    87.23E-01
                     59.69E-01
                                        60.00E-01
                                                            0.050
    86.61E-01
                     64.64E-01
                                        65.00E-01
                                                            0.050
    85.94E-01
                     70.00E-01
                                        70.00E-01
                                                            0.050
                     74.76E-01
    85.41E-01
                                        75.00E-01
                                                            0.050
    84 .90E-01
                     79.71E-01
                                        80.00E-01
                                                            0.050
    84.44E-01
                     84.93E-01
                                        85.00E-01
                                                            0.050
    84.01E-01
                     90.25E-01
                                        90.00E-01
                                                            0.050
```

95.00E-01

10.00E+00

10.50E+00

11.00E+00

11.50E+00

12.00E+00

0.050

0.050

0.050

0.050

0.050

0.050

94.98E-01

10.04E+00

10.46E+00

10.96E+00

11.49E+00

12.01E+00

Time and Date: 11:46:50 10-16-1986 - 12:48:13 10-16-1986

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82.91E-01

82.58E-01

82.28E-01

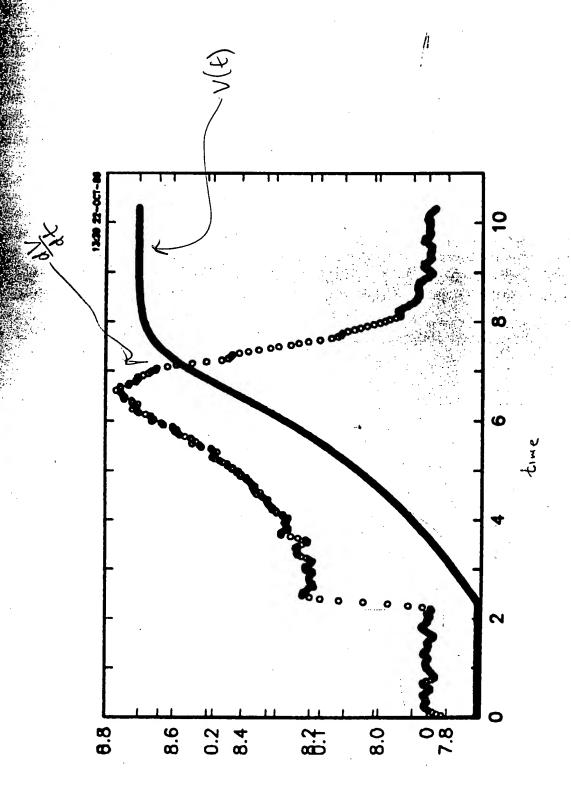
82.01E-01

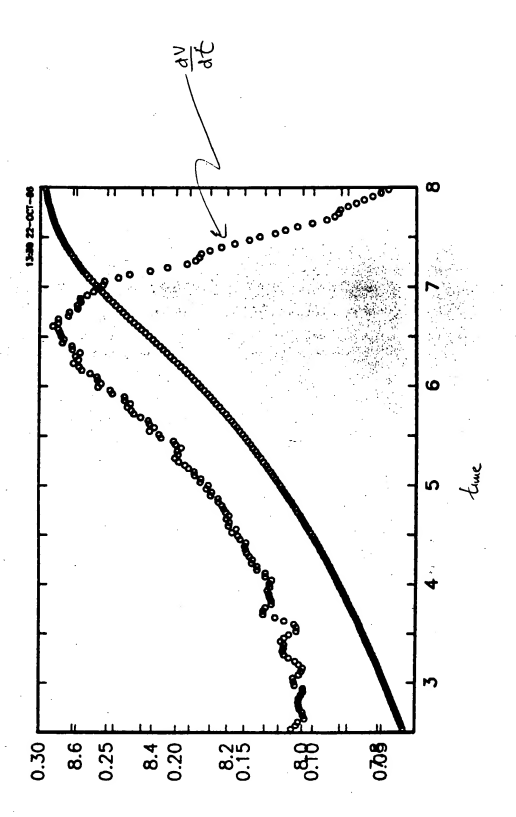
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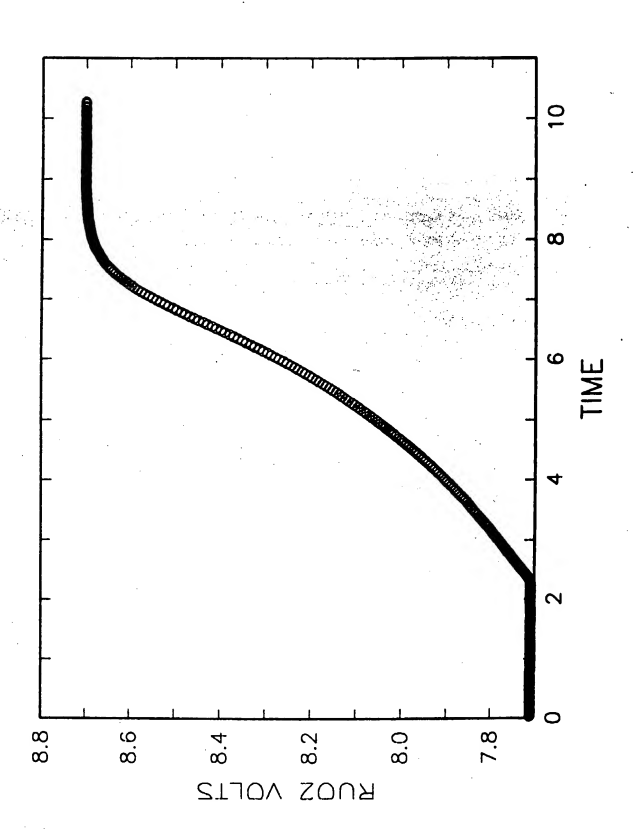
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Min Volts to Plot = 8.00 Volts
Max Volts to Plot = 10.0 Volts
Plot Title
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X Axis Plot Label = Sample Temp (K)
Y Axis Plot Label = RU02 (Volts)
Start Temperature = 002 K
Temperature Step = 0.5 \text{ K}
Temperature Stop = 014 K
LR400 Range
                 = 20000 \text{ Ohms}
Temperature Dev. = 0.050 K
Wait Time = 0010 seconds
Notebook messages = .4
RUO2 Calibration vs C2329 Carbon Glass Resistor
4 WIRE CONFIGURATION FOR BOLOMETER
BACKGROUND MEASUREMENT.
STEPPING UP
Number of points = 21
                         Values per point = 4
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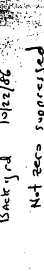
Title: RUO2 VS TEMP PROGRAM

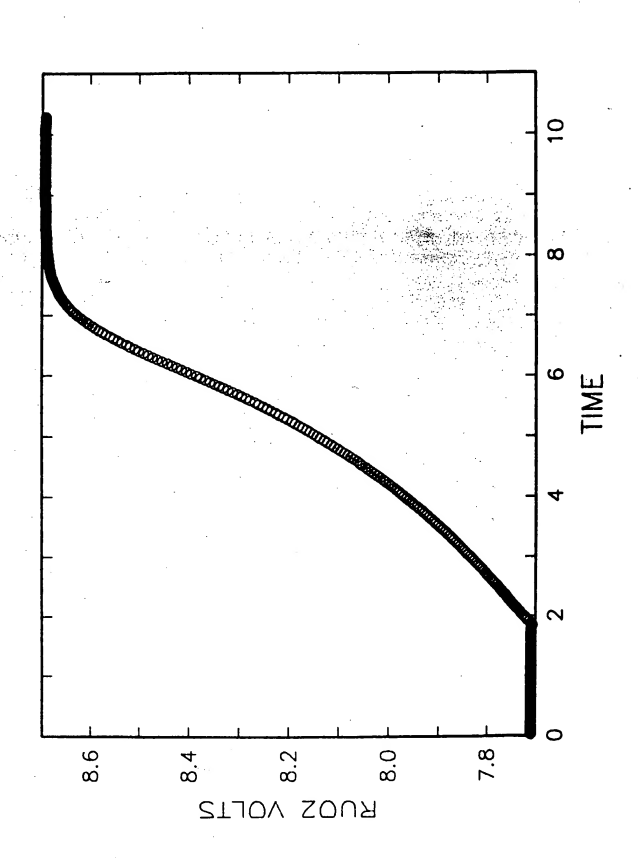
RUO2 VOLTS	C2329 TEMP	SET TEMP	TEMP DEV
99,26E-01	19.96E-01	" 20.00E-01	0.050
96.15E-01			
	25.17E-01	25.00E-01	0.050
94.16E-01	29.68E-01	30.00E-01	0.050
92.41E-01	34.72E-01	35.00E-01	0.050
91.03E-01	39.68E-01	40.00E-01	0.050
89.85E-01	44.67E-01	45.00E-01	0.050
88.85E-01	49.68E-01	50.00E-01	0.050 -
87.98E-01	54.76E-01	55.00E-01	0.050
87.23E-01	59.69E-01	60.00E-01	0.050
86.61E-01	64.64E-01	65.00E-01	0.050
85.94E-01	70.00E-01	70.00E-01	0.050
85.41E-01	74.76E-01	75.00E-01	0.050
84.90E-01	79.71E-01	80.00E-01	0.050
84.44E-01	84.93E-01	85.00E-01	0.050
84.01E-01	90.25E-01	90.00E-01	0.050
83.61E-01	94.98E-01	95.00E-01	0.050
83.25E-01	10.04E+00	10.00E+00	0.050
82.91E-01	10.46E+00	10.50E+00	0.050
82.58E-01	10.96E+00	11.00E+00	0.050
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82.01E-01	12.01E+00	12.00E+00	0.050

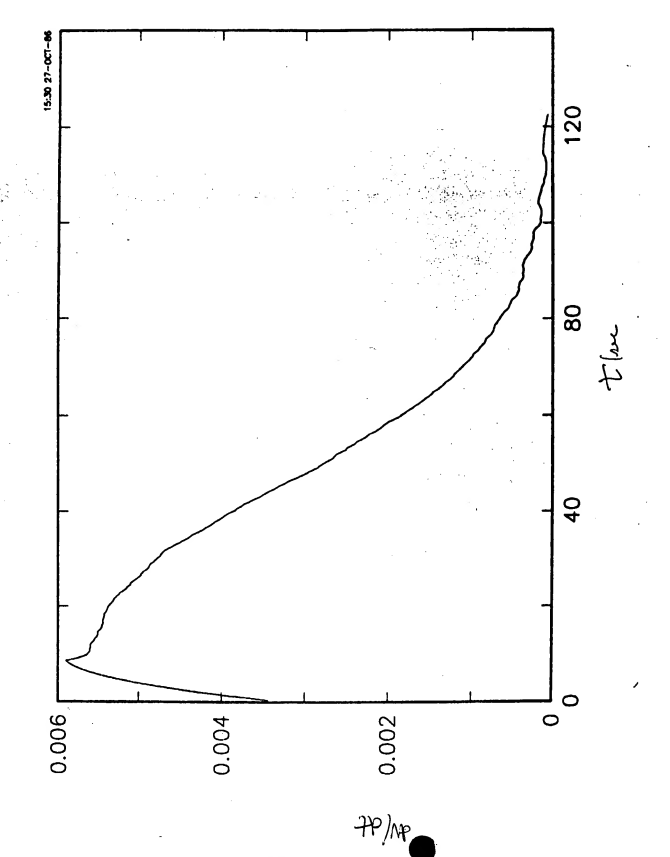


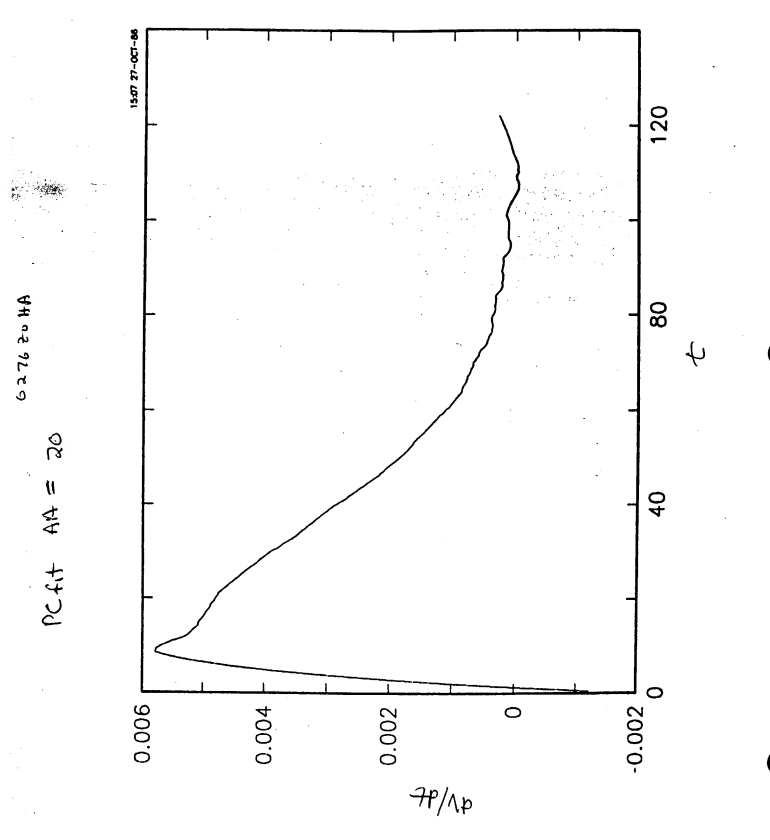


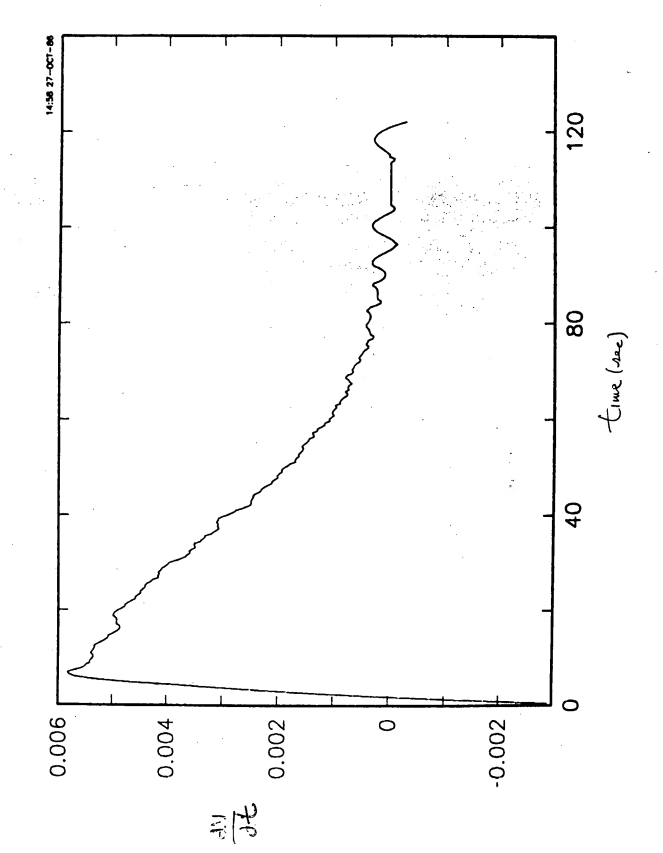


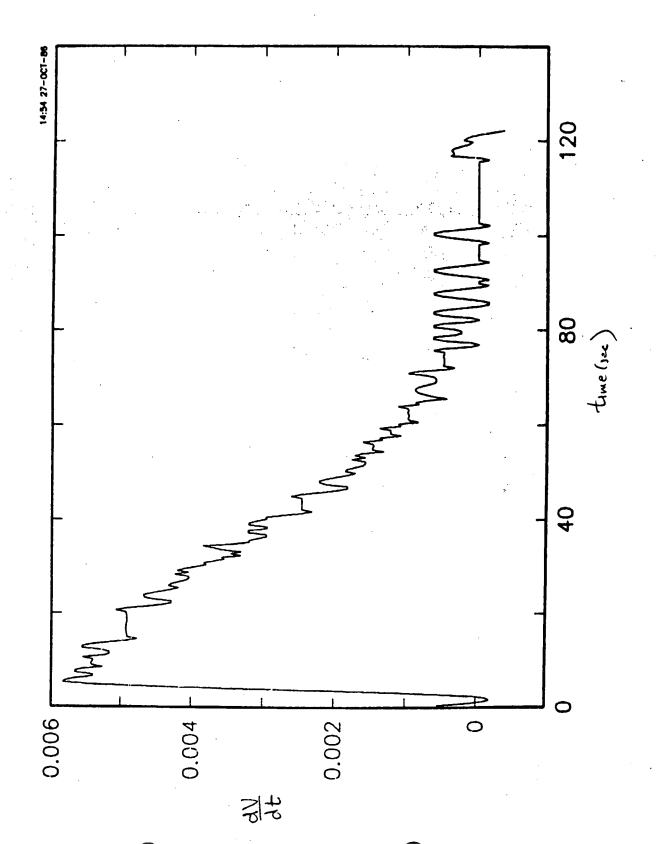


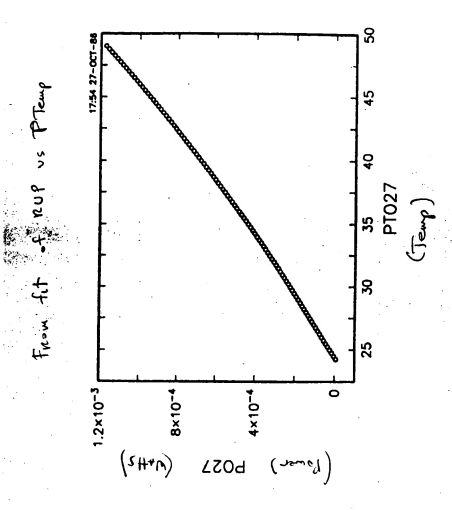


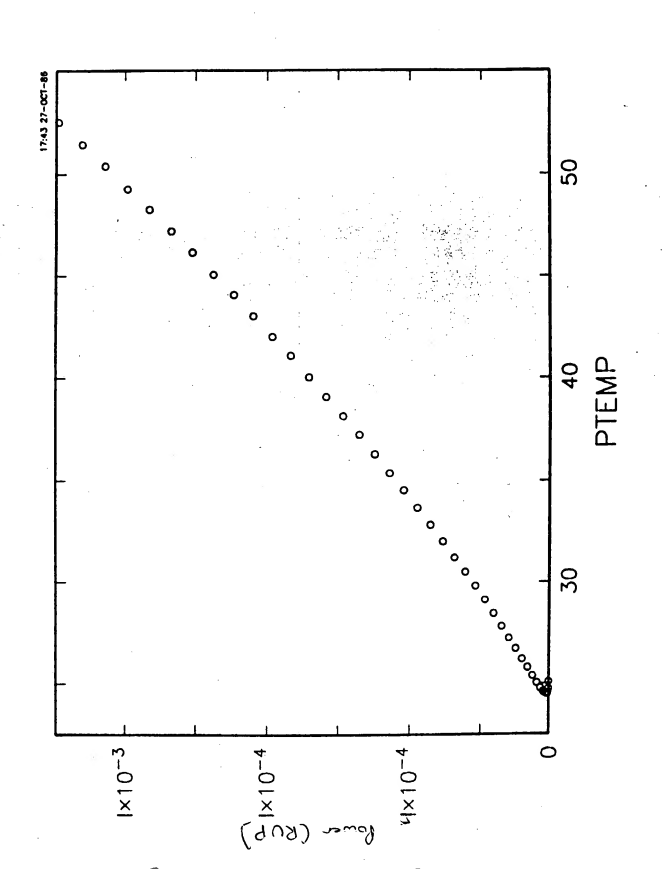


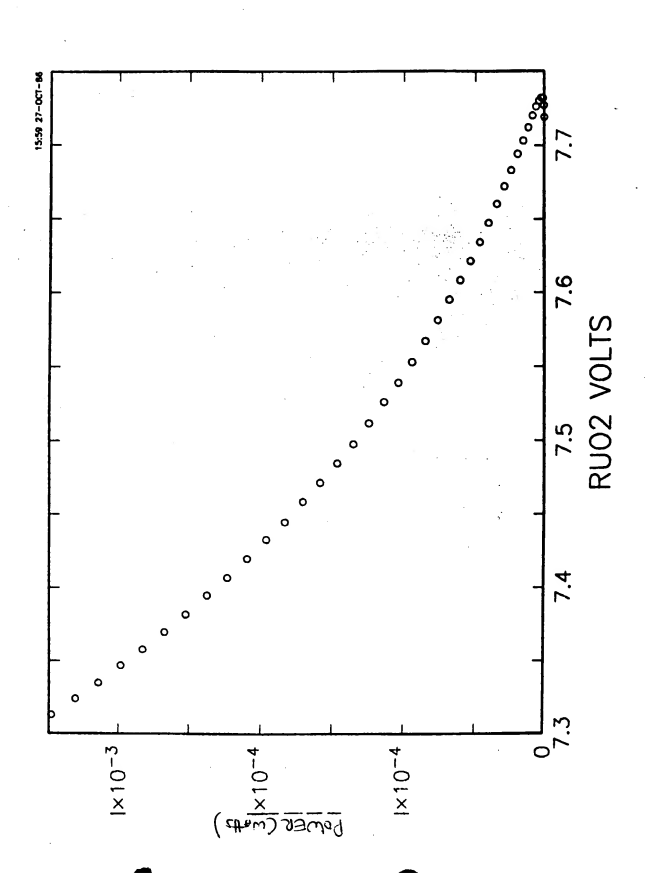




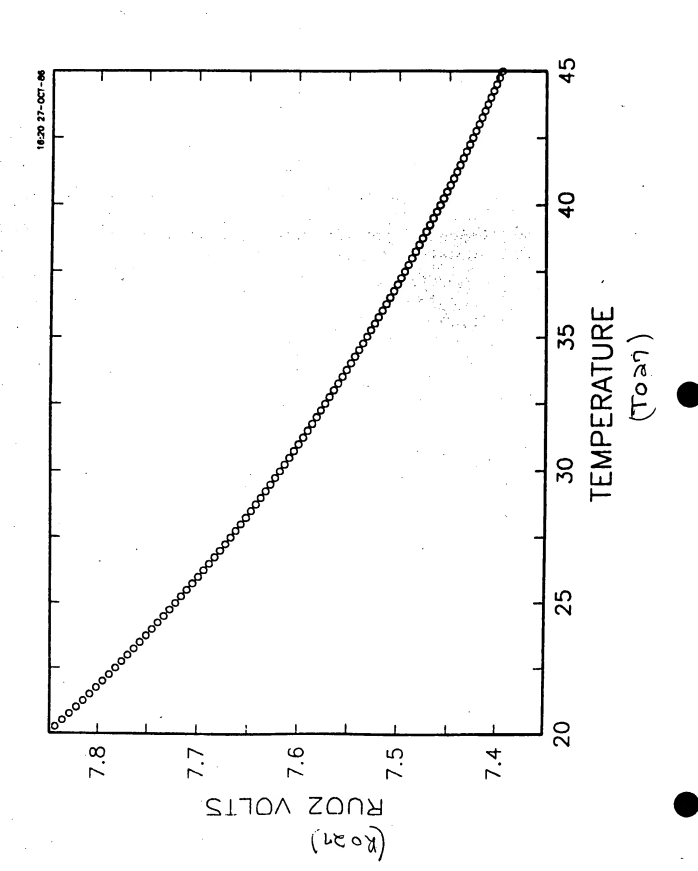


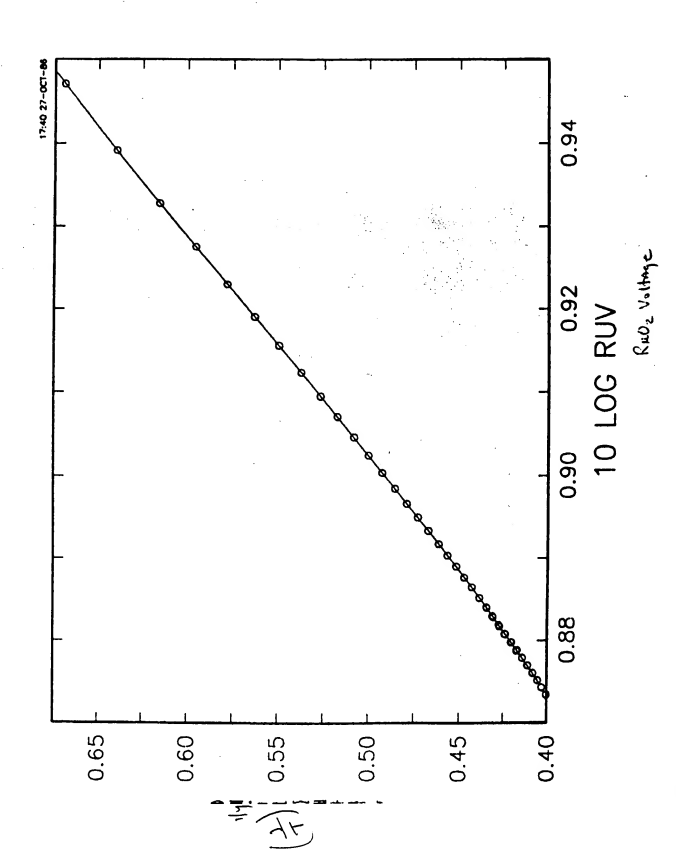


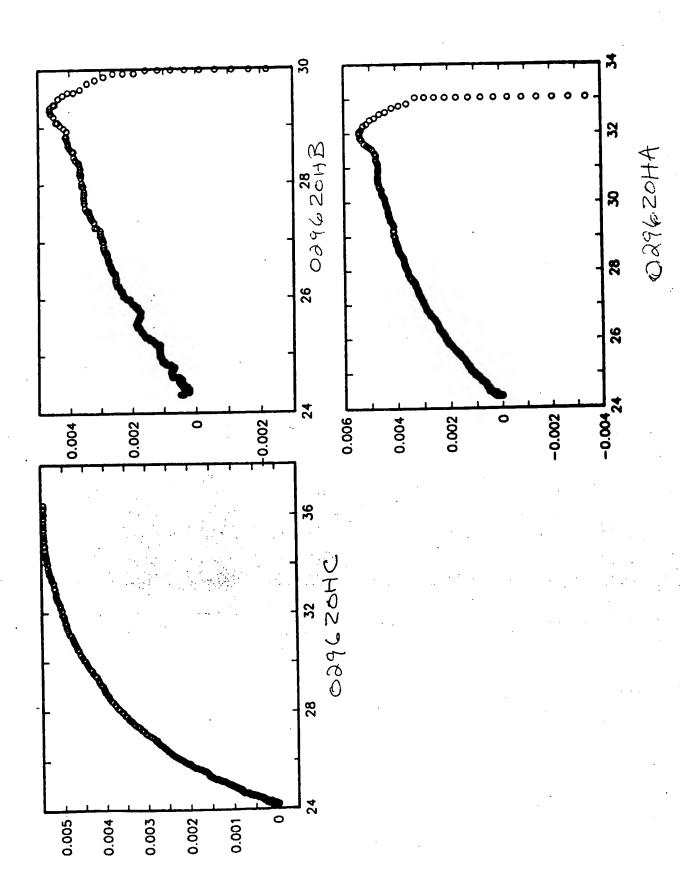


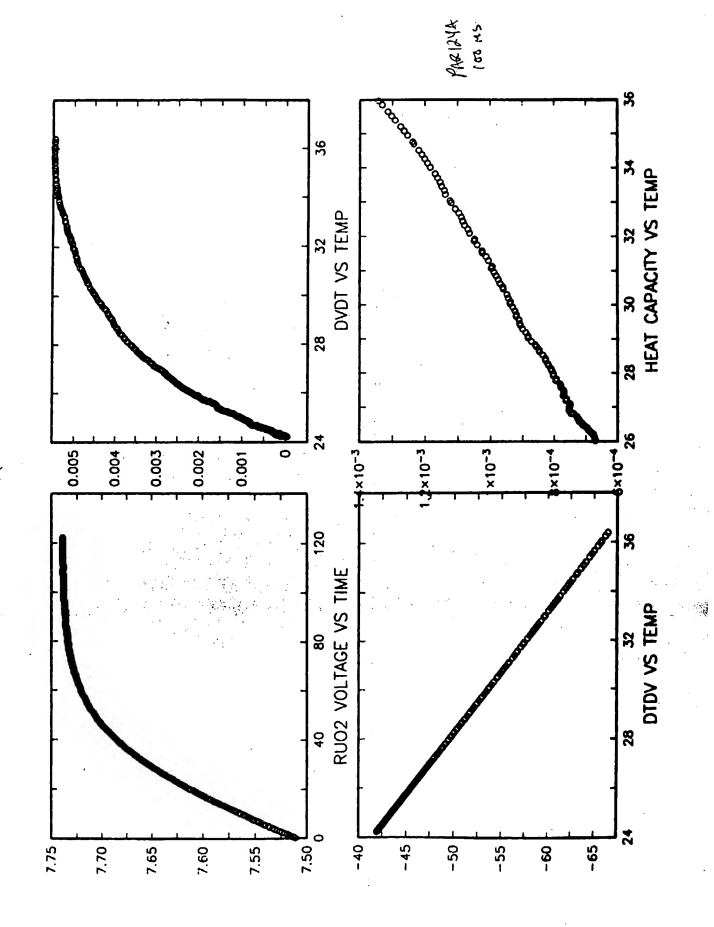


OZIGHTCU.DAT & files for RUOZ CALLEGATION









6] A. VERBAERE and M. TOURNOUX, Bull. Sac. Chim., 1973, 1237.

17] A. D. WADSLEY, Acta Cryst., 1964, 17, 623.

48] H. REDBAII, G. DESGARDIN and B. RAVEAU, J. Solid State Chem., 1980, 31, 321.

19] M. Hervieu, H. Rebani, G. Desgardin and B. Raveau, J. Solid State Chem., 1980, 35, 200.

[20] M. Hervieu and B. Raveau, J. Solid State Chem., 1980, 32, 161.

21) H. Rennah, J. Pannetier and B. Raveau, J. Solid State Chem., 1982, 41, 57, 22) A. Leclaire, J. C. Monier and B. Raveau, Acta Cryst. (to be published).

[記] C. MICHEL, C. ROBERT, D. GROULT and B. RAVEAU, Mal. Res. Bull., 1975, 10,

(Received November 30, 1983).

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in mixed valence copper oxides related Oxygen intercalation to the perovskites

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series of compounds: BasLasCuaO1118, Las-xA1+xCusOa-x/2+8 and Las-xAxCuOa-x/2+8 cies in (001) planes of these structures makes that two of these families: Ba, Ca, Cu, Oit+8 ABSTRACT. — Intercalation of oxygen in ternary copper oxides has been studied for three Cu(II) and Cu(III) siniultanously are oxygen defect compounds whose structure is closely related to that of the perovskite, and to those of the two members of the intergrowths Sro-perovskite: Sr, Ti, O, and K, NiF, respectively. The localization of the oxygen vacanand Laz-xAz+xCuzOz-x/z+8 can be considered in their most reduced state as oxides with (A = Ca, Sr, Ba). These mixed valence copper oxides, characterized by the presence of low dimensionality. The influence of oxygen intercalation on the structure is described. The electrical properties of these phases are described and discussed: they are strongly influenced by the intercalation process. A progressive transition from a p type semiconductive to a p type semi-metallic or metallic state is indeed observed which depends on the oxygen pressure and on the nature of the oxides.

Lag-xAxCuOq-x/2+8(A = Ca, Sr, Ba). Ces oxydes de cuivre à valence mixte, caractérisés dont la structure est étroitement liée respectivement à celle de la pérovskite et à celles des par la présence simultanée de Cu(II) et Cu(III), sont des composés déficitaires en oxygène deux membres de la série d'intercroissances pérovskite-Sro : SrafigO, et KaNiFa. La ocifisation des lacunes anioniques dans les plans (001) de ces structures fait que deux 'intercalation d'oxygene dans la structure est décrite. Les propriétés électriques de ces Résumé. - L'intercalation d'oxygène dans les oxydes ternaires de cuivre a été étudiée de jes samilles : BagLagCuolines et Lag-xAi+xCuolog-xizes peuvent être considérées, dans leur état le plus réduit, comme des oxydes de basse dimensionalité. L'influence de phases sont décrites et discutées : elles sont fortement influencées par le processus d'intercalation. Une transition progressive d'un état semi-conducteur de type p à un état semimétallique ou métallique de même type, qui dépend de la pression d'oxygène et de la pour trois séries de composés : Ba₃La₃Cu₄O₁₄₊₈, La_{1-x}A_{1+x}Cu₄O_{4-x/1+8} nature des oxydes, est en esfet observée.

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Intercalation of oxygen in an oxide, by a simple reversible exchange with O2 in air or in a gascous atmosphere can be used for different applications such as electrocatalysis, or gauges for materials with electrical properties sensitive to the oxygen content. Thus it appears that such oxides must exhibit rather large oxygen defects in their a reduced » form, and must be able to absorb oxygen from atmosphere tending towards a stoichiometric phase in their « oxidized » state. This phenomenon supposes a reversible change of the oxidation state and of the coordination number of the metallic aloms which participate to the framework of the oxide. In this respect, ter oxides are very good candidates, owing to the ability of copper to and several oxidations stades: + 1, + 2, + 3. Cu(11) and Cu(111) must be especially considered owing to their possibility to take the same octahedral take several coordinations—octahedral, square pyramidal, square planar coordination in similar structures as shown from previous works on La2Cu"O4 [1-2] and LaSrCu"O4 [3], which are isostructural with K2NiF4. Ternary oxides ArCuyO3 containing Cu(111) are more difficult to prepare to several kbars [3-8] are most of the time necessary to synthesize these compounds. However, the presence of A elements like barium favours the than those with Cu(II), since oxygen pressures ranging from 1 bar [4-7] formation of Cu(111) in normal pressure conditions [9-10]. The present paper deals with the soft intercalation of oxygen, i. e. at low pressure ($p\leqslant 1$ atm) and at low temperature (T $\sim 400\text{--}500^{\circ}\,\text{C}$) in three series of ternary copper oxides related to the perovskite [11-13] and belonging to the systems La2O3-AO-CúO with A = Ca, Sr, Ba. The influence of oxygen intercalation on the It transport properties of these phases are discussed.

STRUCTURAL CONSIDERATIONS

Three families with an oxygen defect structure have been isolated in the systems La2O3-AO-CuO;

- The oxygen defect perovskites La3Ba3Cu6O14+3.
- The oxygen defect intergrowths Sr3Ti2O, type,
- The oxygen desect intergrowth K2NiF4 lype, La2-xAxCuO4-x/2+6. $La_{2-x}A_{1+x}Cu_2O_{6-x/2+\delta} \Lambda = Ca$, Sr.

The most reduced form which has been isolated for the defect perovskites La, Ba, Cu, Oi4+, corresponds to the formulation La, Ba, Cu, Oi4. Its

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OXYGEN INTERCALATION IN COPPER OXIDES

structure (fig. 1) can be described as an ordered oxygen defect perovskite. All the metallic sites corresponding to the stoichiometric perovskite are occupied by copper ions and lanthanum and barium ions respectively, whereas only 7/9 of the anionic sites are occupied in an ordered manner.

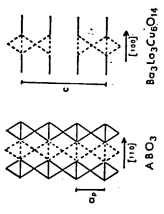


Fig. 1. - Schematic structure of a stoichiometric ABO, perovskite ind the defect oxygen perovskite Ba, La, Cu, Oit.

 $c \simeq 3a_p = 11.721 \text{ A}$), it can indeed be seen that the basal planes of the Considering the tetragonal cell of this compound ($a \simeq a_p \sqrt{2} = 5.525 \text{ Å}$, octaliedra, parallel to (001) are preserved, that one apex out of two is missing at the levels z = 1/6 and 5/6, whereas all the apices of these octahedra are missing at z = 1/2. It results that this reduced form can be considered as a true layer structure: double defect perovskite layers Bal., Lao., 5Cu, O, 1, built up from corner-sharing, octahedra CuO, square pyramids CuO, and square groups CuO, are observed whose cohesion is ensured by lanthanum ions located at z = 1/2. It is remarkable that such an oxide is characterized by a high Cu(111) content in spite of the high oxygen desect content. Site potential calculations confirm that the Cu3+ ions are located preferentially on the octahedral sites. It must also be noted that this at 1000° C for 24 h the mixture of La2O3, CuO and BaCO3 and quenching the samples at room-temperature a slight excess of oxygen is indeed to bserved phasi La, Ba, Cu, O14.05 is then synthesized by annealing the sample limit compound has not really been synthesized. By heating in air corresponding to the formulation La3Ba3Cu6O14.10. The most reduced La, Ba, Cu, O14.10 at 400° C under low oxygen pressure (~ 5.10-3 bar) during several hours.

The deviation from stoichiometry in the oxides La2-xA1+xCu2O6-x/2+3 lium for lanthanum, in a small homogeneity range ($0 \le x \le 0.14$ for is more complex owing to the possibility of substitution of calcium or stron-

strontium and x=0.10 for calcium). The most reduced oxide which has been isolated in this family corresponds to the formulation La₂SrCu₂O₆. Its tetragonal cell (a = 3.865 Å, c = 19.887 Å), corresponds to a structure closely related to that of $Sr_3Ti_2O_3$ (fig. 2 a). Cu^{2+} ions are indeed located on the Ti++ sites. La3+ and Sr2+ ions are located on the Sr2+ sites, whereas thus, this oxide can be considered as an intergrowth of double oxygen six anionic sites out of seven are occupied by oxygen in an ordered manner; perovskite layers and SrO type layers. The perovskite layers exhibit some similarity with those ebserved for La, Ba, Cu,O14: the basal planes of the octahedra parallel to (0.0.1) are also preserved whereas at z=0 and z = 1/2 all the apices of the oxygen octahedra are missing. However, the resulting configuration of the framework is different from La $_3$ Ba $_3$ Cu $_6$ O $_{_14}$: rertheless this oxide, like La3Ba3Cu6O14, must be considered as a struc-(11) exhibits here only one coordination which is square pyramidal. ture with low dimensionality. It can indeed be described as built up from Sr^{2+} and La^{3+} ions located at z=0 and z=1/2. The | LaSrCu₂O₆ |_m slabs | LaSrCu2On | parallel to (001) whose cohesion is ensured by slabs are themselves an intergrowth of SrO-type layers and corner-sharing structure: the latter corresponds indeed to the superposition of two square pyramid layers. Such slabs are in fact derived from the K2NiF4 | K2Ni2F6 | slabs which would share the face of their square pyramids, forming NiF, octahedra (fig. 2 h). Like LajBajCu,O14+s, LazSrCu2O, is

Statizo, tozs. Cuzo, Kznir, Kz

Fig. 2.

Schematic structure of SraTiO; and LasSrCusO, (projection on to (100) plane),

Schematic representation of K, Ni, Fe slabs sharing the square faces of the NiF, pyramids

characterized by a great stability in spite of its oxygen defect structure: it is indeed synthesized by heating the stoichiometric mixture of CuO, La₂O₃ and SrCO₃ at 1 050-1 100° for 24 h in air and by queryching them at room temperature in order to avoid their oxidation at lower temperature. Contrary to La₃Ba₃Cu₆O₁₄, copper is in its lower oxidation state, Cu(II) in this oxide.

The oxides $La_{2-x}A_xCuO_{4-x/2+\delta}$ exhibit an oxygen defect K_2NiF_4 type structure involving different coordinations of copper: octahedral, square pyramidal and eventually square planar (fig. 3). Their oxygen content

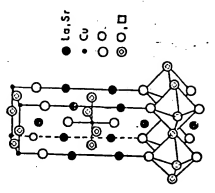
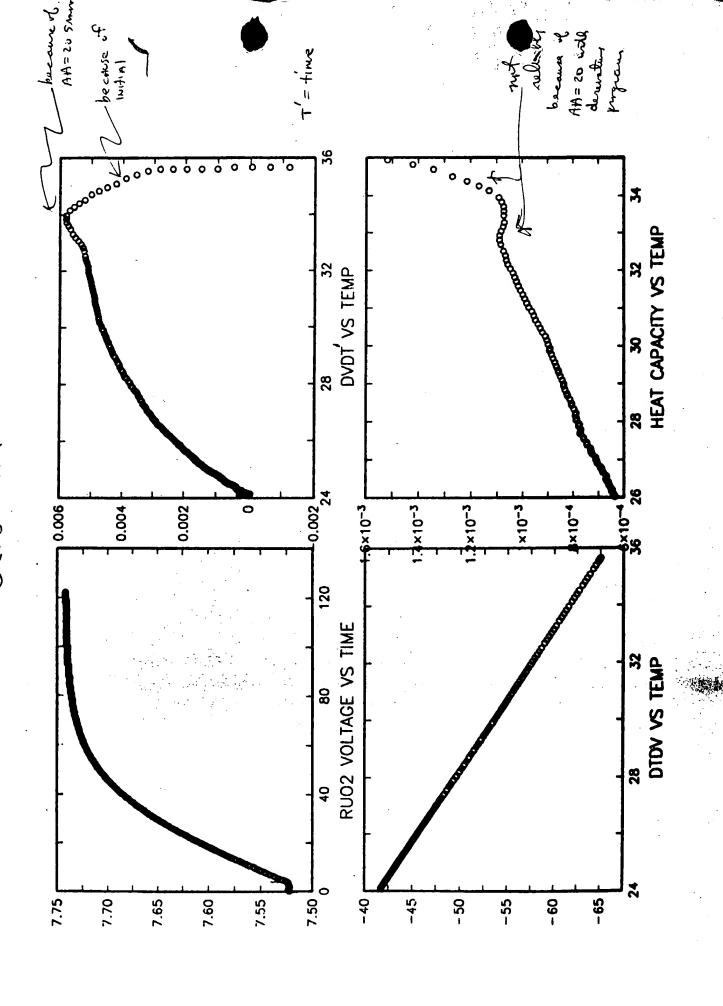


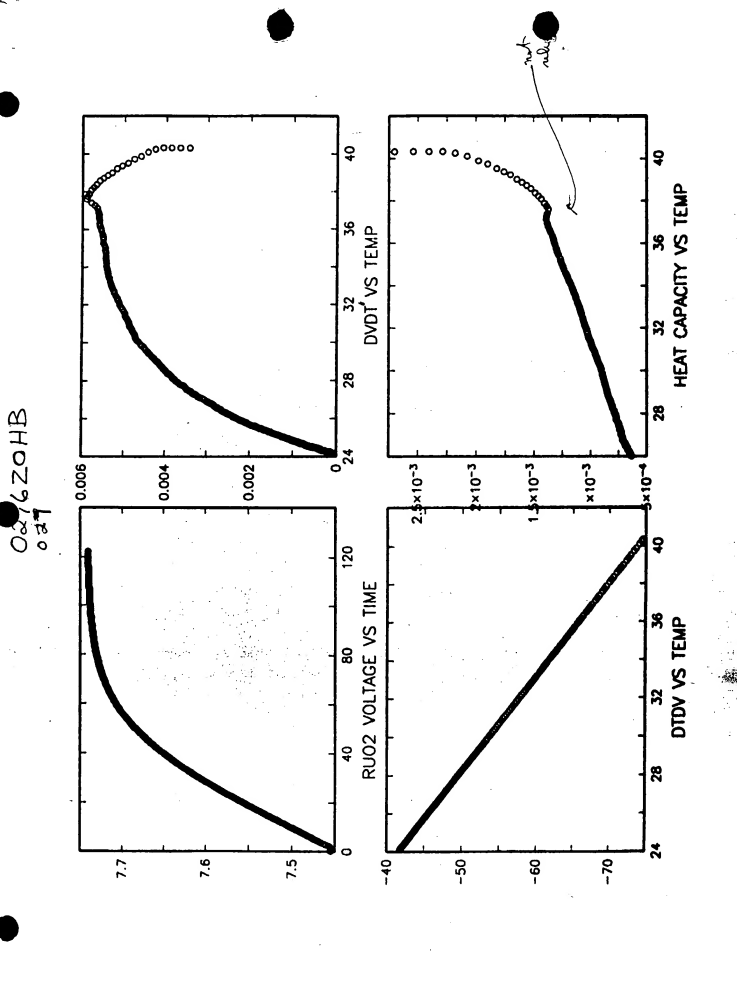
Fig. 3. — Perspective view of the structure of the oxides $\text{La}_{1-x}\text{Sr}_x\text{CuO}_{4-x/3+8}$ with oxygen vacancies located in the basal plane of the octahedra.

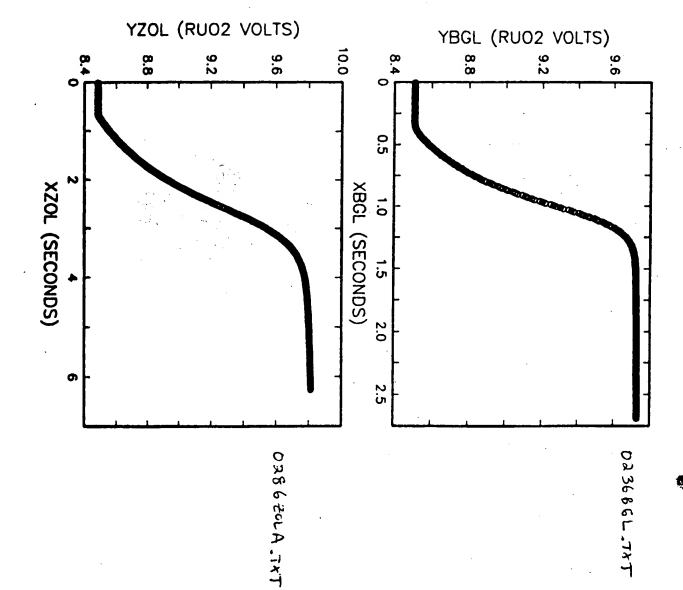
depends on the nature of the A ions ($\Lambda = \text{Ca}_1^3 \text{Sr}$, Ba) and on the substitution rate x which can lead to wide homogeneity ranges: $0 \le x \le 0.20$ for $\Lambda = \text{Ca}$ and Ba and $0 \le x \le 4/3$ for $\Lambda = \text{Sr}$. The most reduced phase which exhibits the highest deviation from stoichiometry has been synthesized in the case of strontium for x = 4/3: La_{2/3}Sr_{4/3}CuO_{3.33}. Contrary to the two other series, the oxygen vacancies are located in the basal plane of the octahedra which are parallel to the (00 1) plane of the tetragopal cell (a = 3.759 Å, c = 12.907 Å). It must also be emphasized that this type of localization of the oxygen vacancies is always observed whatever the nature of the A ions, and whatever the rate of substitution x may be. However, symmetry changes and order-disorder phenomena in this plane may appear according to the nature of A and x value (table 1). So, the calcium and barium oxides are characterized by a monoclinic distortion of the tetragonal K₂NiF₄ structure, whatever the x value may be $0 \le x \le 0.20$; the same

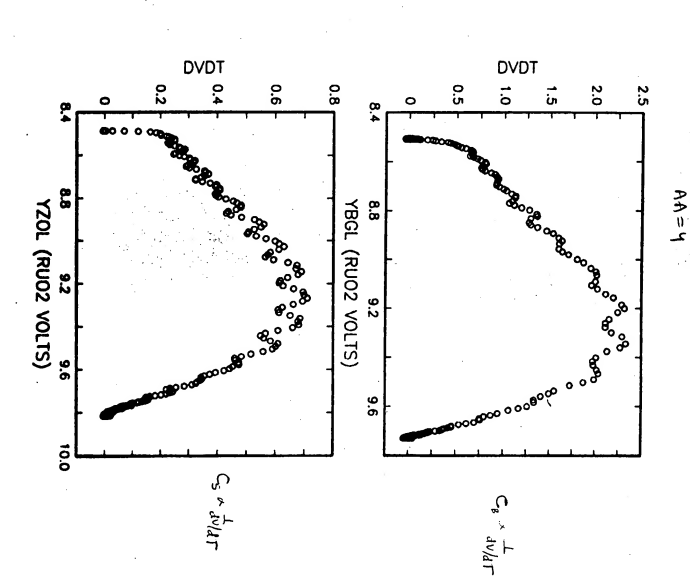
REVUE DE CHIMIE MINÉRALE

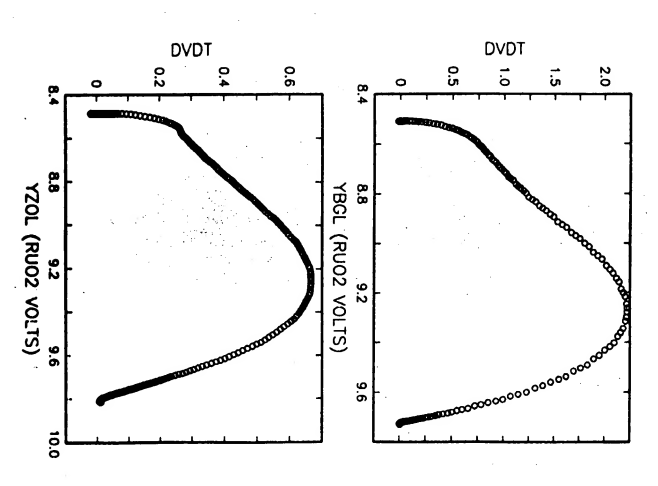
O27620HA

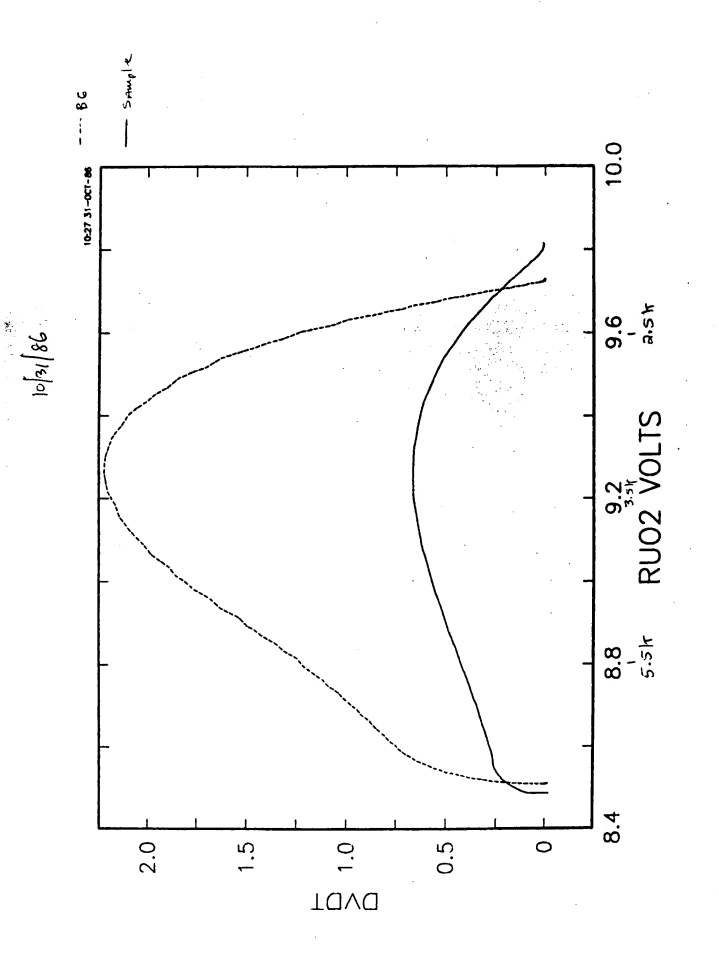


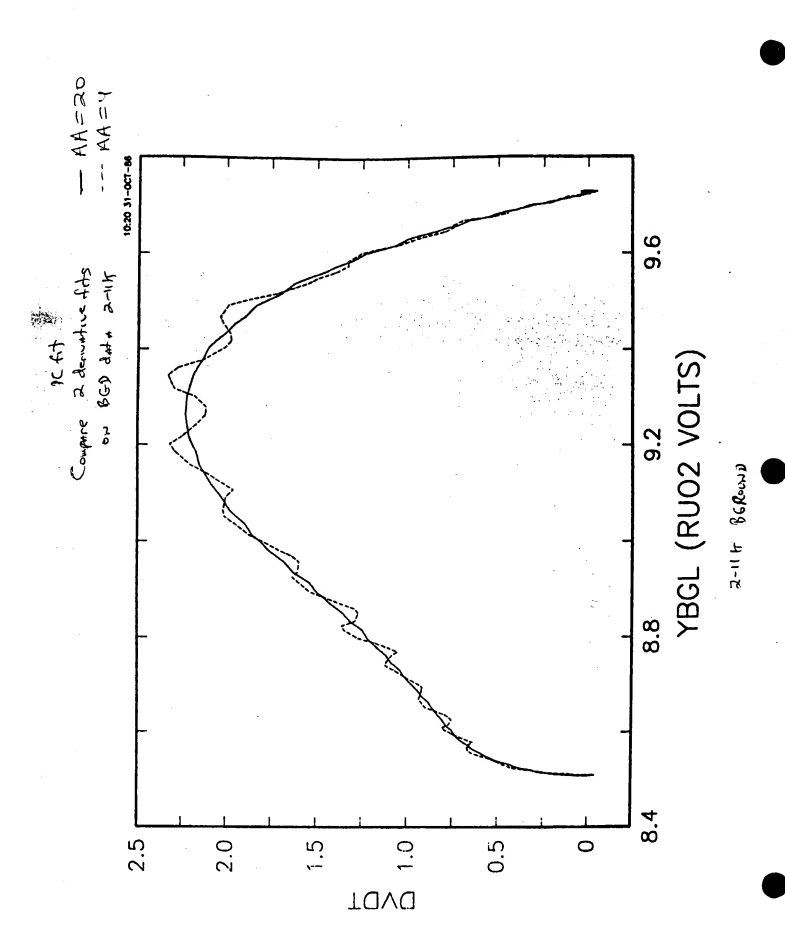


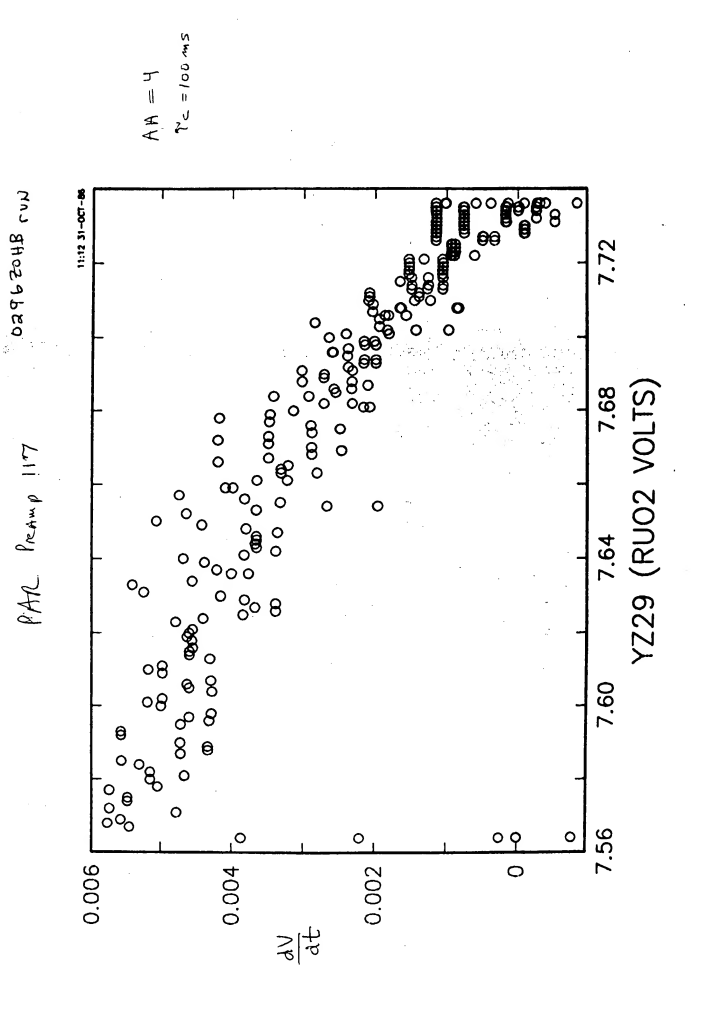


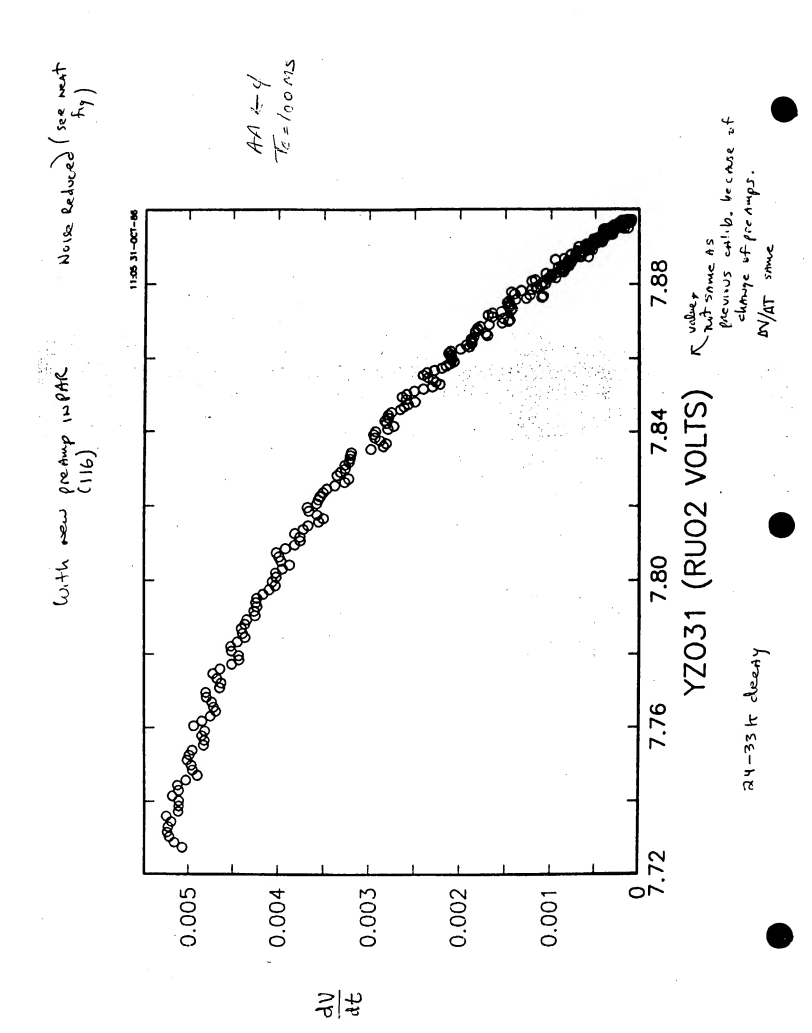












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Min Volts to Plot = 7.48 Volts
Max Volts to Plot = 7.88
                        Volts
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X Axis Plot Label = Sample Temp (K)
Y Axis Plot Label = RU02 (Volts)
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Temperature Step = 1.0 K
Temperature Stop = 040 K
LR400 Range
                 = 00200 Ohms
Temperature Dev. = 0.100 K
Wait Time
              = 0015 seconds
Notebook messages = 4
RUO2 Calibration vs C2329 Carbon Glass Resistor
4 WIRE CONFIGURATION FOR BOLOMETER
TEMP. CALIBRATION 26-40K
STEPPING UP
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                          Values per point = 4
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78.21E-01	28.08E+00	28.00E+00	0.100
78.11E-01	29.07E+00	29.00E+00	0.100
77.95E-01	30.07E+00	30.00E+00	0.100
77.76E-01	31.09E+00	31.00E+00	0.100
77.58E-01	32.09E+00	32.00E+00	0.100
77.41E-01	33.06E+00	33.00E+00	0.100
77.20E-01	34.01E+00	34.00E+00	0.100
77.06E-01	35.06E+00	35.00E+00	0.100
76.90E-01	36.10E+00	36.00E+00	0.100
76.75E-01	37.06E+00	37.00E+00	0.100
76.63E-01	37.92E+00	38.00E+00	0.100
76.45E-01	39.09E+00	39.00E+00	0.100

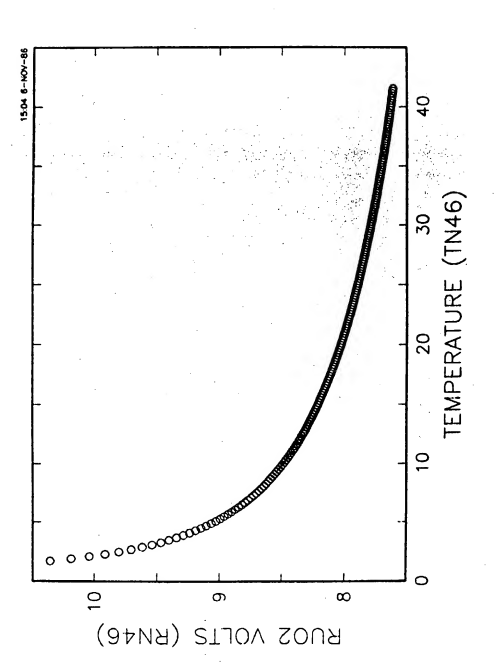
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Max Temp to Plot = 32.00 K
Min Volts to Flot = 7.58 Volts
Max Volts to Plot = 9.11 Volts
Plot Title
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X Axis Plot Label = Sample Temp (K)
Y Axis Plot Label = RUO2 (Volts)
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Temperature Step = 1.0 K
Temperature Stop = 032 K
                  = 02000 \text{ Ohms}
LR400 Range
Temperature Dev.
                  = 0.050 K
Wait Time
              = 0010 seconds
Notebook messages = . 4
RUO2 Calibration vs C2329 Carbon Glass Resistor
4 WIRE CONFIGURATION FOR BOLOMETER
TEMP. CALIBRATION 05-32K
STEPPING UP
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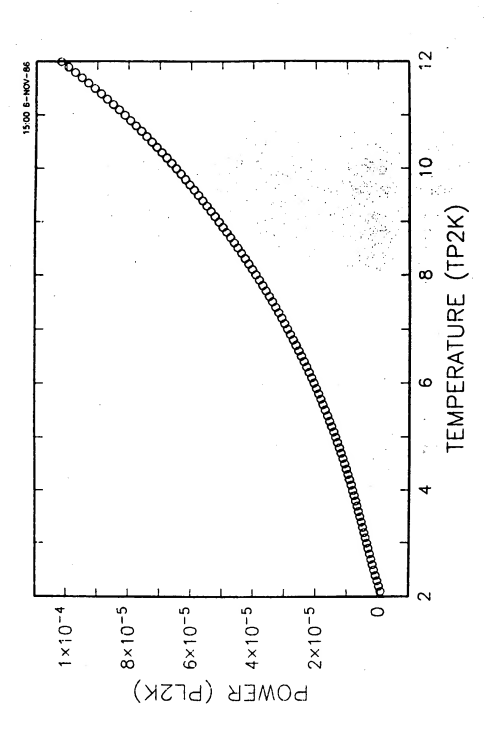
•				
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	0.050	60.00E-01	59.68E-01	88.95E-01
	0.050	70.00E-01	69.80E-01	87.66E-01
	0.050	80.00E-01	79.74E-01	86.61E-01
	0.050	90.00E-01	89.73E-01	85.71E-01
	0.050	10.00E+00	99.81E-01	84.93E-01
. *	0.050	11.00E+00	10.99E+00	84.25E-01
	0.050	12.00E+00	12.00E+00	83.66E-01
	0.050	13.00E+00	13.00E+00	83.13E-01
Section 1	0.050	14.00E+00	13.99E+00	82.67E-01
	0.050	15.00E+00	15.02E+00	82.17E-01
	0.050	16.00E+00	16.00E+00	81.72E-01
	0.050	17.00E+00	17.00E+00	81.33E-01
•	0.050	18.00E+00	18.01E+00	80.96E-01
	0.050	19.00E+00	19.00E+00	80.62E-01
	0.050	20.00E+00	20.02E+00	80.27E-01
	0.050	21.00E+00	20.99E+00	79.99E-01
	0.050	22.00E+00	21.96E+00	79.71E-01
	0.050	23.00E+00	22.98E+00	79.44E-01
	0.050	24.00E+00	23.99E+00	79.18E-01
	0.050	25.00E+00	25.01E+00	78.94E-01
	0.050	26.00E+00	25.99E+00	78.71E-01
- had point (too high for	0.050	27.00E+00	27.03E+00	→78.65E-01
had point (too high for Ruoz volts)	0.050 =	28.00E+00	28.03E+00	78.29E-01
	0.050	29.00E+00	28.99E+00	78.06E-01
	0.050	30.00E+00	29.97E+00	77.86E-01
	0.050	31.00E+00	30.98E+00	77.67E-01

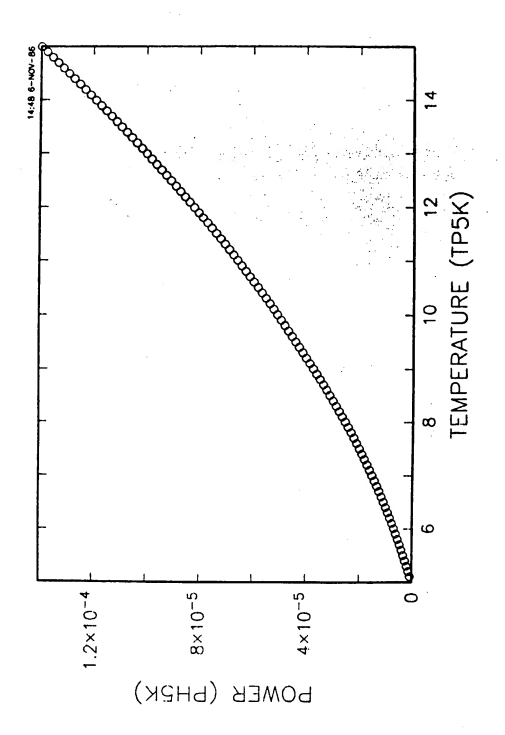
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Max Volts to Plot = 10.4 Volts
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Y Axis Plot Label = RUO2 (Volts)
Start Temperature = 1.7 K
Temperature Step = 0.5 \text{ K}
Temperature Stop = 011 K
LR400 Range
             = 20000 Ohms
Temperature Dev. = 0.050 \text{ K}
Wait Time = 0010 seconds
Notebook messages = 4
RUO2 Calibration vs C2329 Carbon Glass Resistor
4 WIRE CONFIGURATION FOR BOLOMETER
TEMP. CALIBRATION 1.7-11K
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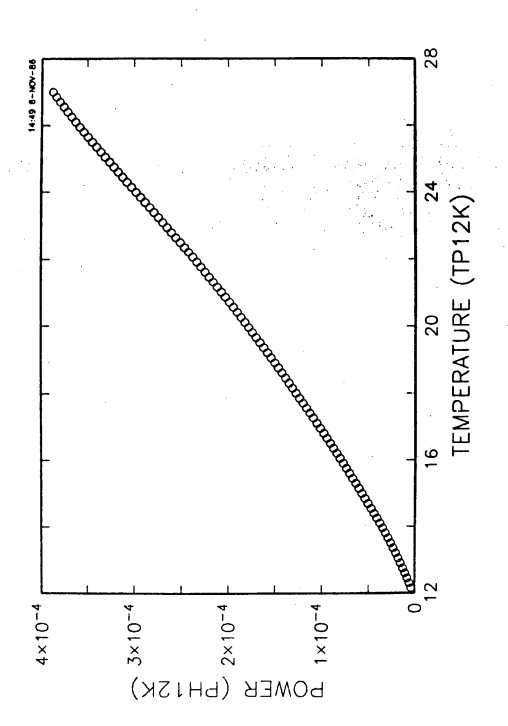
STEPPING UP
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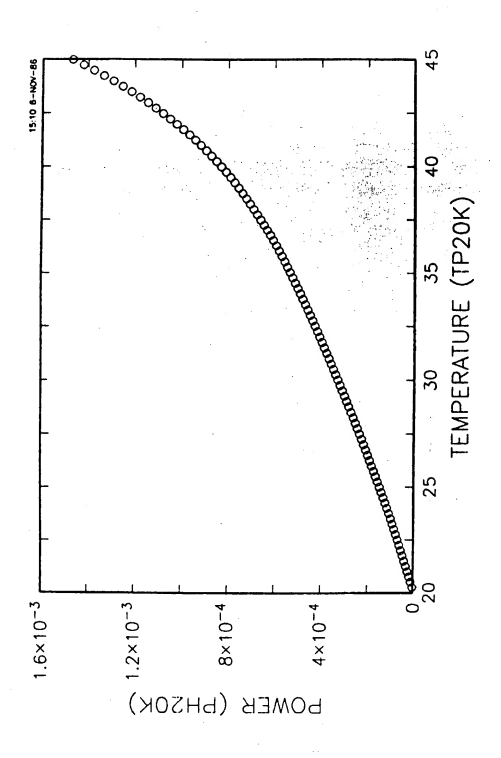
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97.09E-01	26.84E-01	27.00E-01	0.050
95.05E-01	31.84E-01	32.00E-01	0.050
93.42E-01	36.96E-01	37.00E-01	0.050
92.11E-01	42.05E-01	42.00E-01	0.050
91.21E-01	46.96E-01	47.00E-01	0.050
90.14E-01	52.04E-01	52.00E-01	0.050
89.30E-01	57.19E-01	57.00E-01	0.050
88.57E-01	62.17E-01	62.00E-01	0.050
87.92E-01	67.16E-01	67.00E-01	• 0.050
87.32E-01	71.94E-01	72.00E-01	0.050
86.79E-01	76.82E-01	77.00E-01	0.050
86.31E-01	81.74E-01	82.00E-01	0.050
85.85E-01	86.80E-01	87.00E-01	0.050
85.44E-01	91.90E-01	92.00E-01	0.050
85.03E-01	97.25E-01	97.00E-01	0.050
84.70E-01	10.23E+00	10.20E+00	0.050

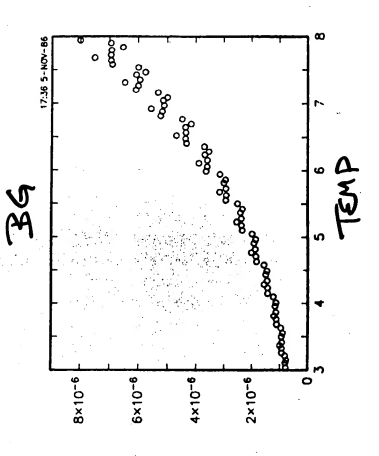




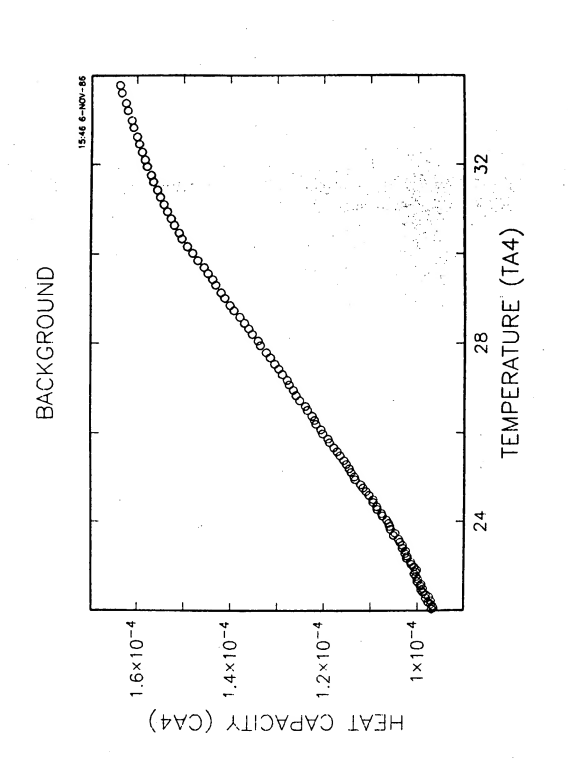


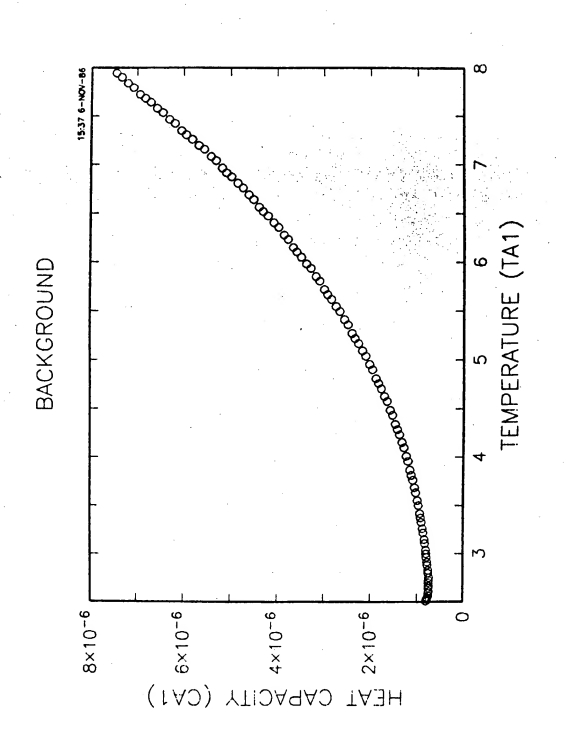


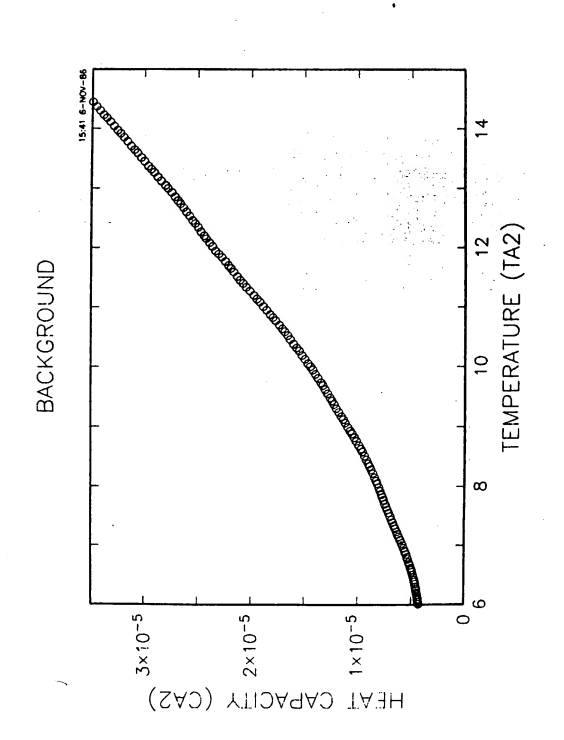


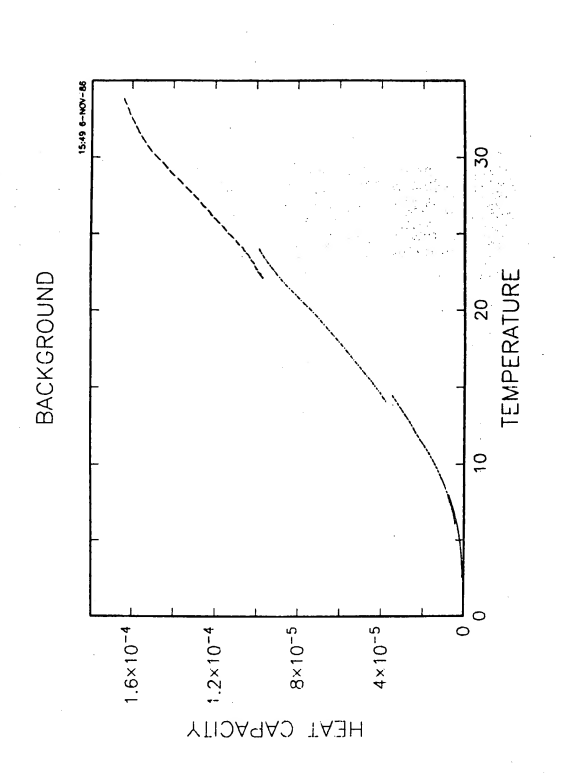


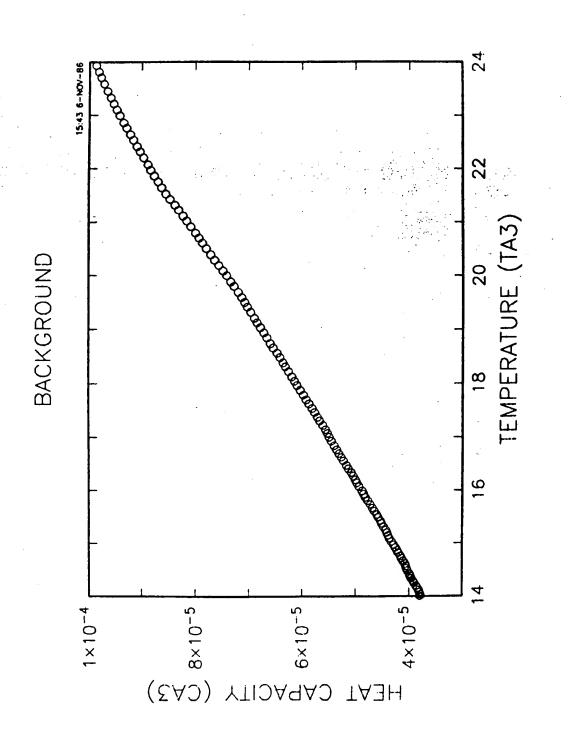
BACKGROUND

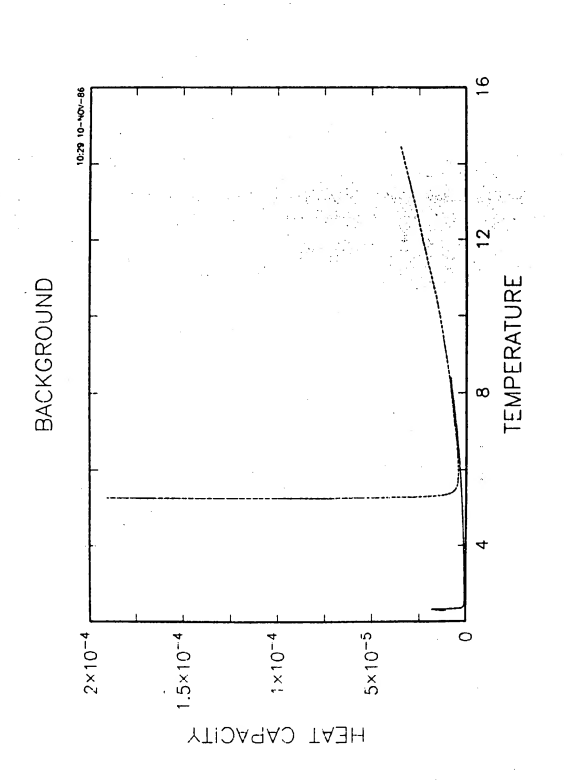


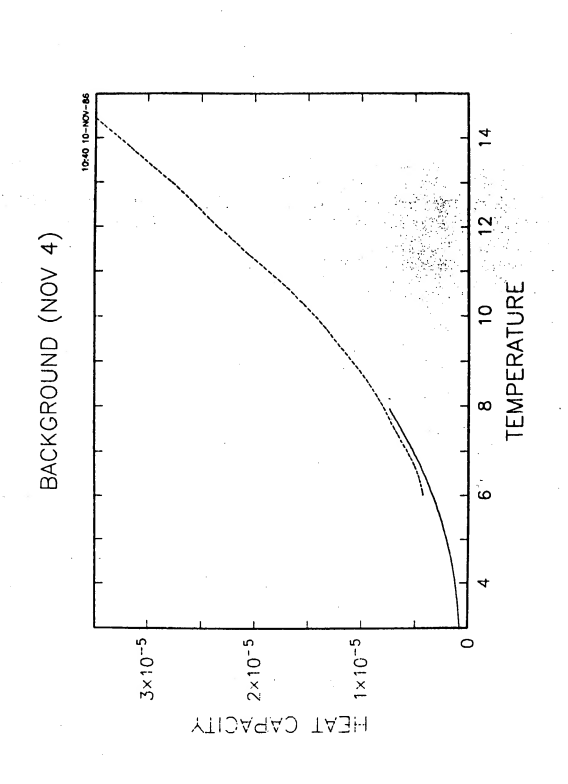


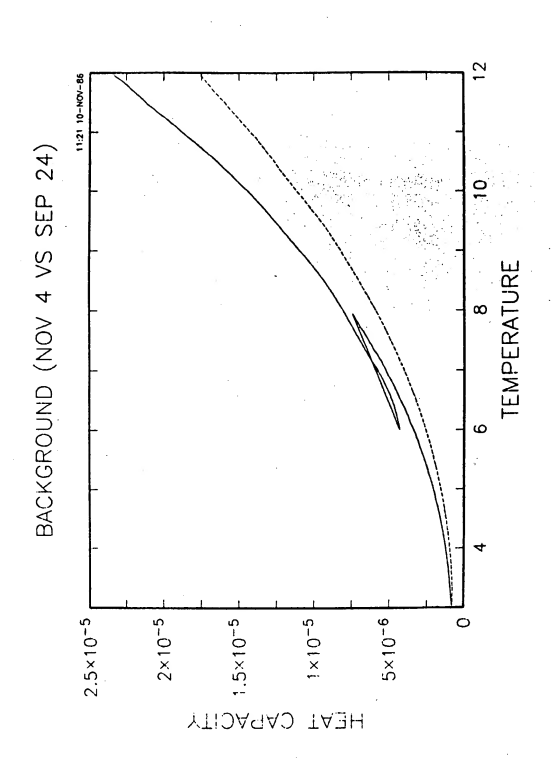


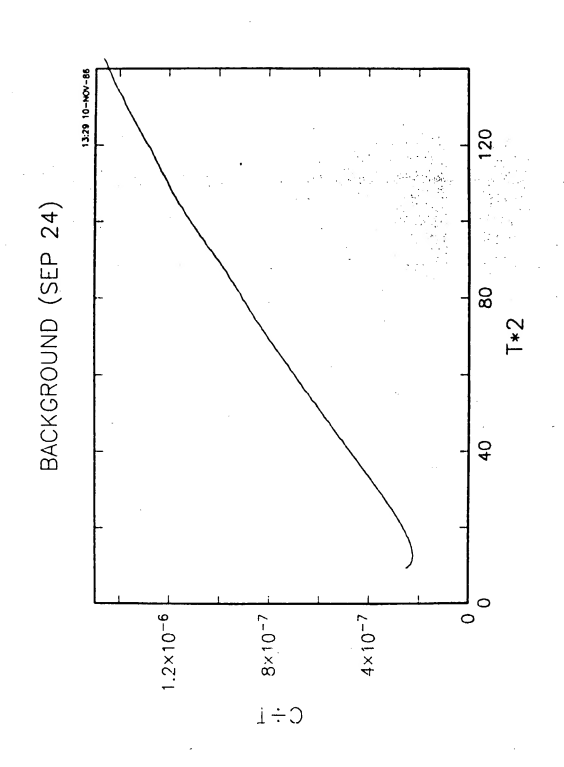


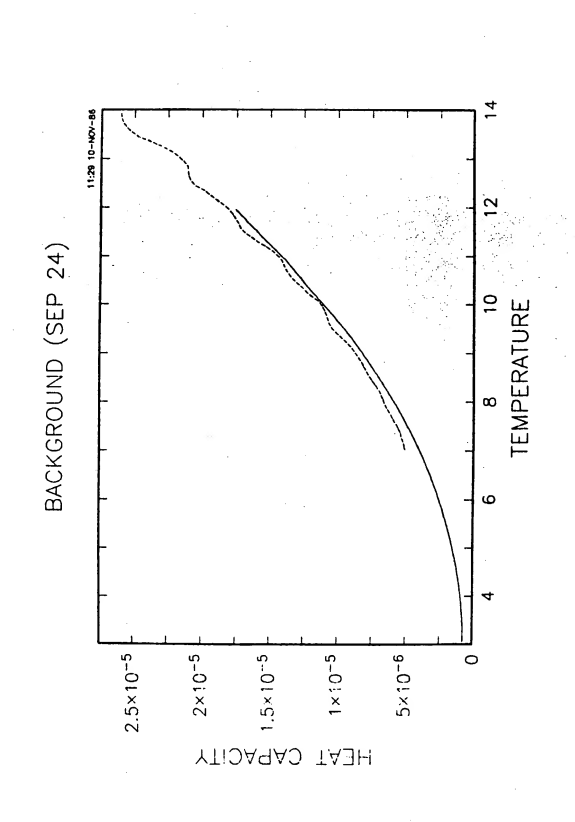


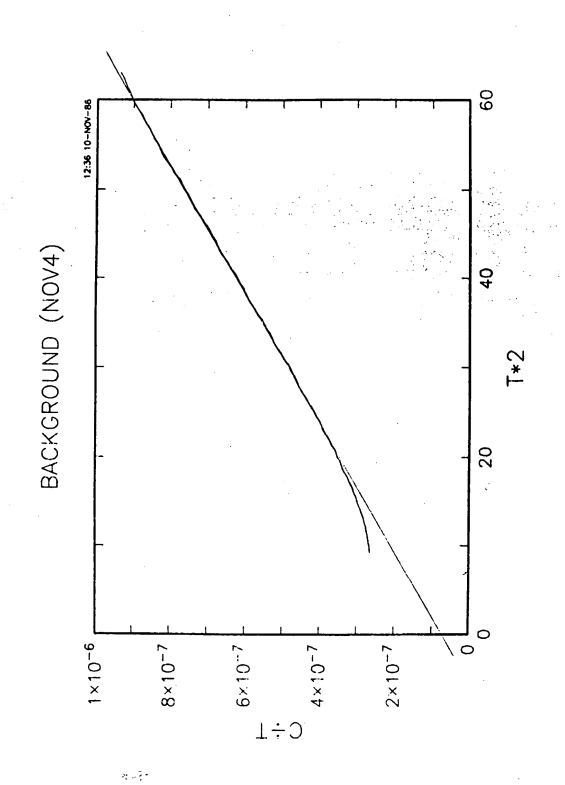


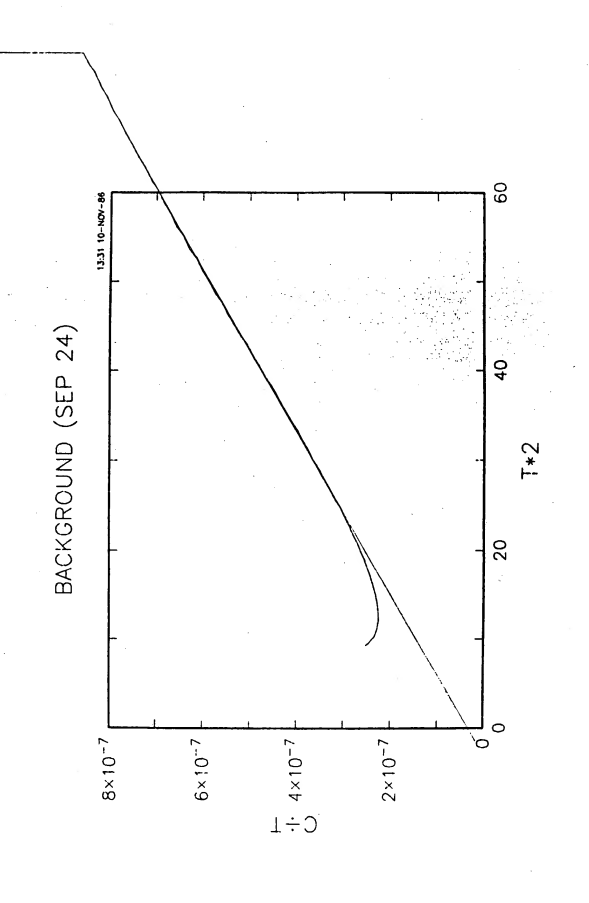


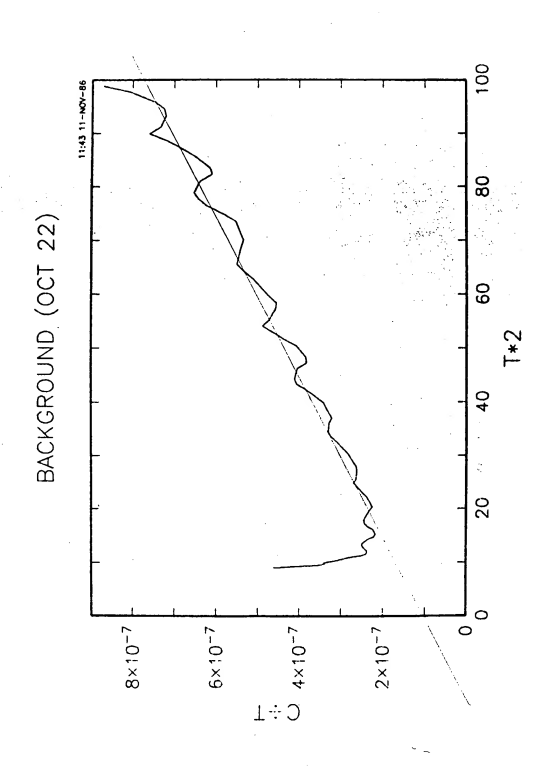


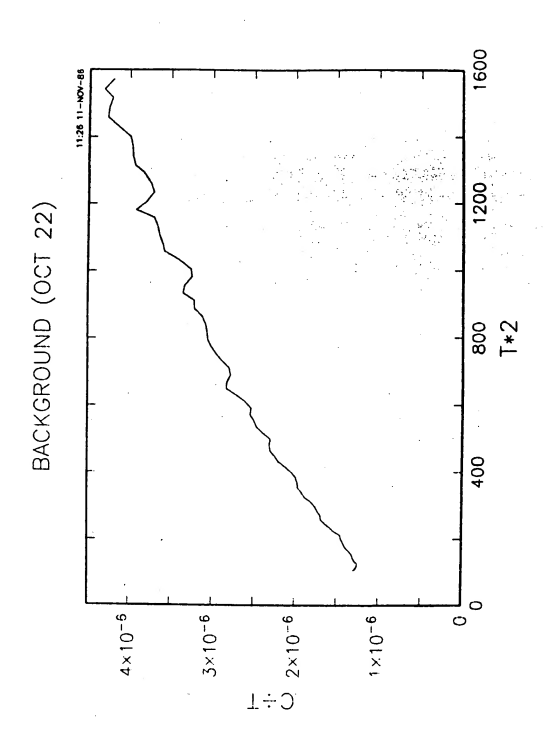












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Title: RUO2 VS TEMP PROGRAM
Run Id. - NO76LTCA.DAT
 ime and Date: 07:35:26 11-03-1986 - 08:11:58 11-93-1986
  rameter lines = 13
Min Temp to Plot = 1.500 K
Max Temp to Plot = 12.50 \text{ K}
Min Volts to Plot = 8.30 Volts
Max Volts to Plot = 10.5 Volts
Plot Title
                  = RUO2 VS TEMP
X Axis Plot Label = Sample Temp (K)
Y Axis Plot Label = RU02 (Volts)
Start Temperature = 002 K
Temperature Step = 0.5 \text{ K}
Temperature Stop = 010 K
LR400 Range
                  = 20000 Ohms
Temperature Dev. = 0.050 \text{ K}
Wait Time = 0010 seconds
Notebook messages = 4
RUO2 Calibration vs C2329 Carbon Glass Resistor
4 WIRE CONFIGURATION FOR BOLOMETER
TEMP. CALIBRATION 2-12K
STEPPING UP
Number of points = 15
                          Values per point = 4
```

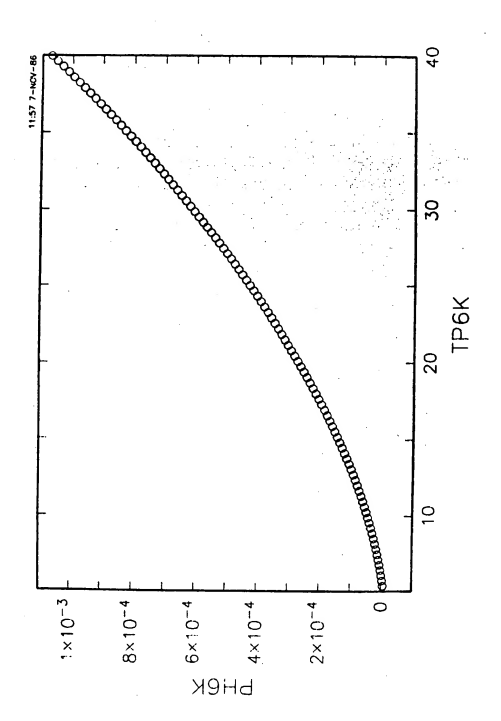
	,		
RUO2 VOLTS	C2329 TEMP	SET TEMP	TEMP DEV
10.18E+00	19.85E-01	20.00E-01	0.050
98.68E-01	24.86E-01	25.00E-01	0.050
96.38E-01	29.87E-01	30.00E-01	0.050
94.60E-01	34.94E-01	35.00E-01	0.050
93.29E-01	39.99E-01	40.00E-01	0.050
92.07E-01	45.01E-01	45.00E-01	0.050
91.02E-01	49.97E-01	50.00E-01	0.050
90.20E-01	54.65E-01	55.00E-01	0.050
89.37E-01	60.15E-01	60.00E-01	0.050
88.69E-01	45.33E-01	45.00E-01	0.050
88.09E-01	70.19E-01	70.00E-01	0.050
87.54E-01	75.08E-01	75.00E-01	0.050
87.04E-01	80.30E-01	80.00E-01	0.050
86.60E-01	85.18E-01	85.00E-01	0.050
86.12E-01	90.30E-01	90.00E-01	0.050

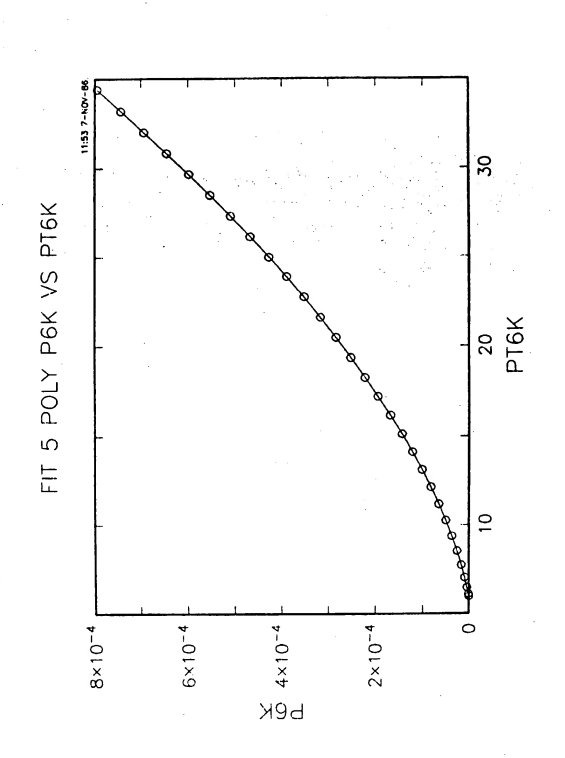
Title: RUO2 VS TEMP PROGRAM Run Id. - NOSSHTCA.DAT Time and Date: 07:56:24 11-02-1986 - 10:29:36 11-42-1986 rameter lines = 13Min Temp to Plot = 05.00 K Max Temp to Plot = 40.00 KMin Volts to Plot = 7.50 Volts Max Volts to Plot = 9.10 Volts Plot Title = RUO2 VS TEMP X Axis Plot Label = Sample Temp (K) Y Axis Plot Label = RUO2 (Volts) Start Temperature = 5.0 K Temperature Step = 1.0 K Temperature Stop = 040 K LR400 Range = 02000 Ohms Temperature Dev. = 0.100 K Wait Time = 0012 seconds Notebook messages = 4 RUO2 Calibration vs C2329 Carbon Glass Resistor 4 WIRE CONFIGURATION FOR BOLOMETER TEMP. CALIBRATION 5-40K STEPPING UP Number of points = 26 Values per point = 4

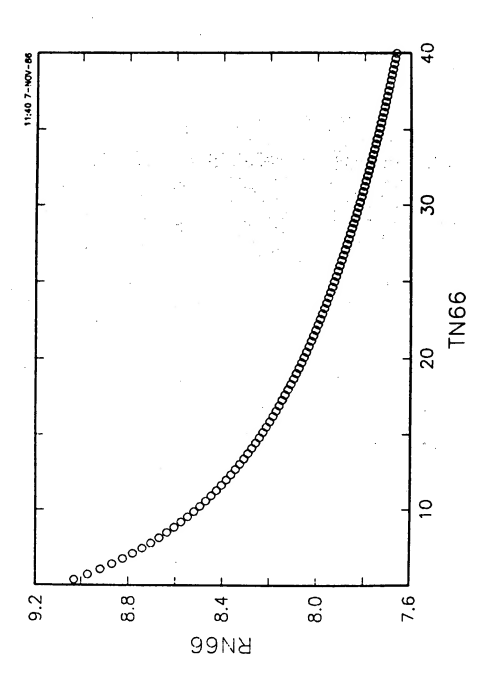
RUO2 VOLTS	C2329 TEMP	SET TEMP	TEMP DELL
91.04E-01	49.67E-01	50.00E-01	TEMP DEV
89.37E-01	59.54E-01	60.00E-01	0.100
88.03E-01	69.63E-01	70.00E-01	0.100
86.85E-01	79.56E-01	80.00E-01	0.100
85.95E-01	89.48E-01	90.00E-01	0.100
85.17E-01	99.64E-01	10.00E+00	0.100
84.49E-01	10.96E+00		0.100
83.87E-01	12.00E+00	11.00E+00	0.100
83.33E-01		12.00E+00	0.100
82.84E-01	12.99E+00	13.00E+00	0.100
	13.98E+00	14.00E+00	0.100
82.38E-01	14.98E+00	15.00E+00	0.100 -
81.96E-01	15.99E+00	16.00E+00	0.100
81.56E-01	17.00E+00	17.00E+00	0.100
81.20E-01	18.00E+00	18.00E+00	0.100
80.87E-01	19.03E+00	19.00E+00	0.100
80.57E-01	19.96E+00	20.00E+00	0.100
80.28E-01	20.96E+00	21.00E+00	0.100
80.02E-01	21.99E+00	22.00E+00	0.100
79.75E-01	23.00E+00	23.00E+00	0.100
79.50E-01	23.96E+00	24.00E+00	0.100
79.27E-01	24.99E+00	25.00E+00	0.100
79.04E-01	25.93E+00	24.00E+00	0.100
78.82E-01	26.99E+00	27.00E+00	0.100
78.62E-01	28.07E+00	28.00E+00	0.100
78.41E-01	28.94E+00	29.00E+00	0.100
78.22E-01	30.02E+00	30.00E+00	0.100
			OFICO

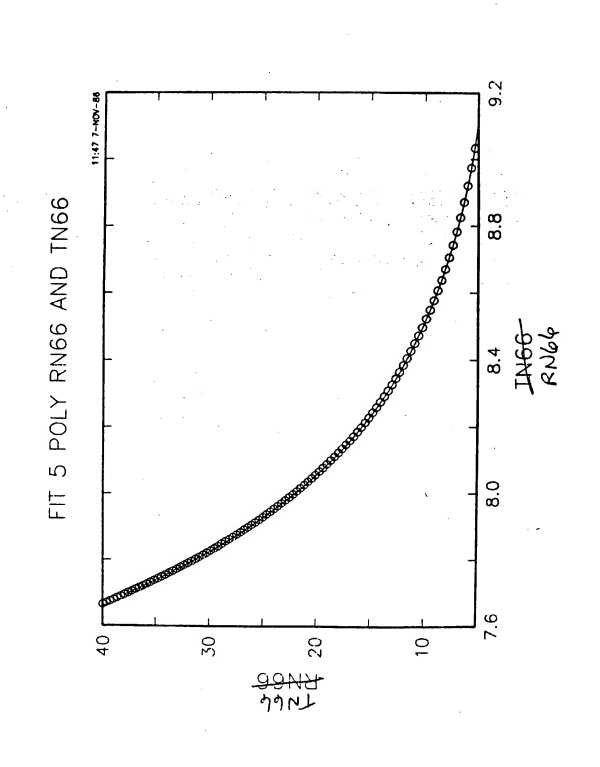
```
Title: RUO2 VS TEMP PROGRAM
 ⊇n Id. - NOS6HTCB.DAT
 me and Date: 10:37:35 11-02-1986 - 11:24:09 11-02-1986
Parameter lines = 13
Min Temp to Plot = 25.00 K
Max Temp to Plot = 40.00 K
Min Volts to Plot = 7.50 Volts
Max Volts to Plot = 7.95 Volts
Plot Title
                 = RUO2 VS TEMP
X Axis Plot Label = Sample Temp (K)
Y Axis Plot Label = RUO2 (Volts)
Start Temperature = 026 K
Temperature Step = 1.0 K
Temperature Stop = 040 K
LR400 Range
                = 02000 Ohms
Temperature Dev. = 0.200 K
Wait Time = 0012 seconds
Notebook messages = 4
RUO2 Calibration vs C2329 Carbon Glass Resistor
4 WIRE CONFIGURATION FOR BOLOMETER
TEMP. CALIBRATION 26-40K
STEPPING UP
                         Values per point = 4
Number of points = 14
```

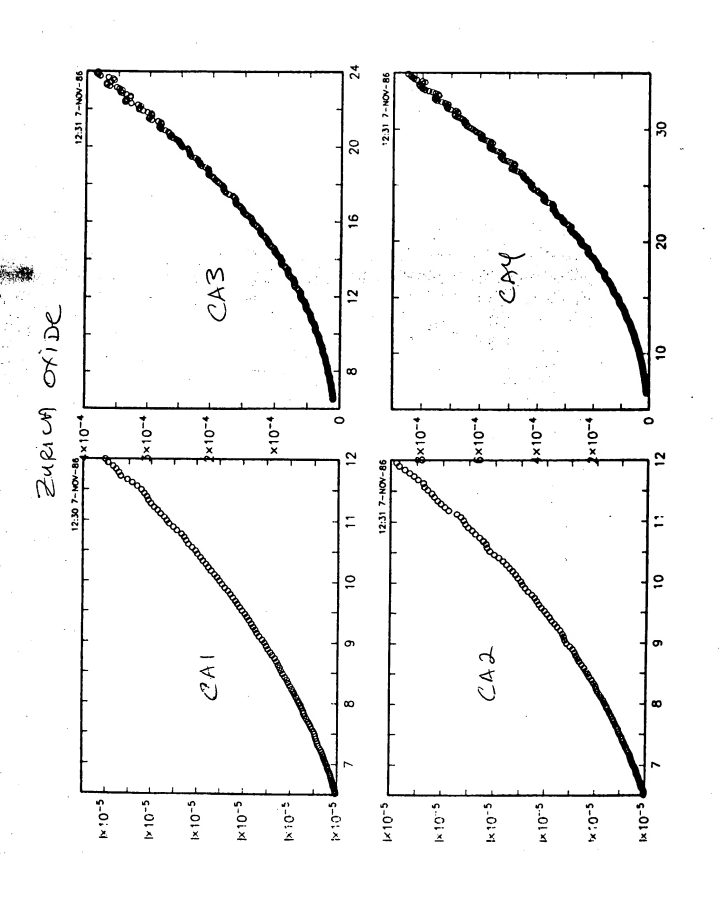
	RUO2 VOLTS	C2329 TEMP	SET TEMP	TEMP DEV
	79.08E-01	25.82E+00	24.00E+00	0.200
_	78.87E-01	27.13E+00	27.00E+00	0.200
	78.64E-01	28.18E+00	28.00E+00	0.200
	78.44E-01	29.09E+00	29.00E+00	0.200
	78.25E-01	30.13E+00	30.00E+00	0.200
	78.06E-01	31.14E+00	31.00E+00	0.200
	77.88E-01	32.13E+00	32.00E+00	0.200
	77.72E-01	33.03E+00	33.00E+00	0.200
	77.55E-01	33.96E+00	34.00E+00	- 0.200
	77.39E-01	35.02E+00	35.00E+00	0.200
	77.22E-01	36.09E+00	36.00E+00	0.200
	77.06E-01	37.03E+00	37.00E+00	0.200
	76.91E-01	38.06E+00	38.00E+00	0.200
	76.75E-01	39.15E+00	39.00E+00	0.200

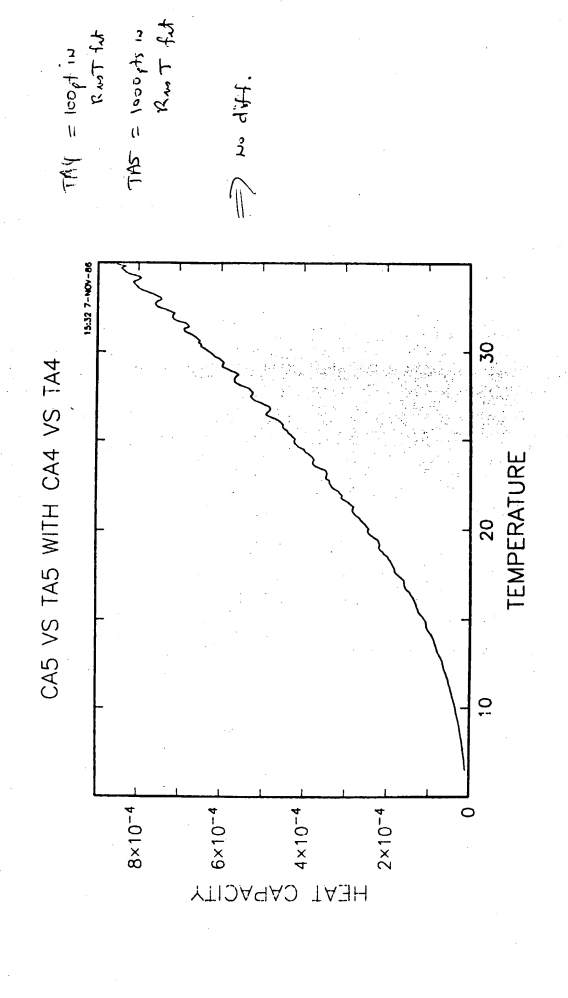


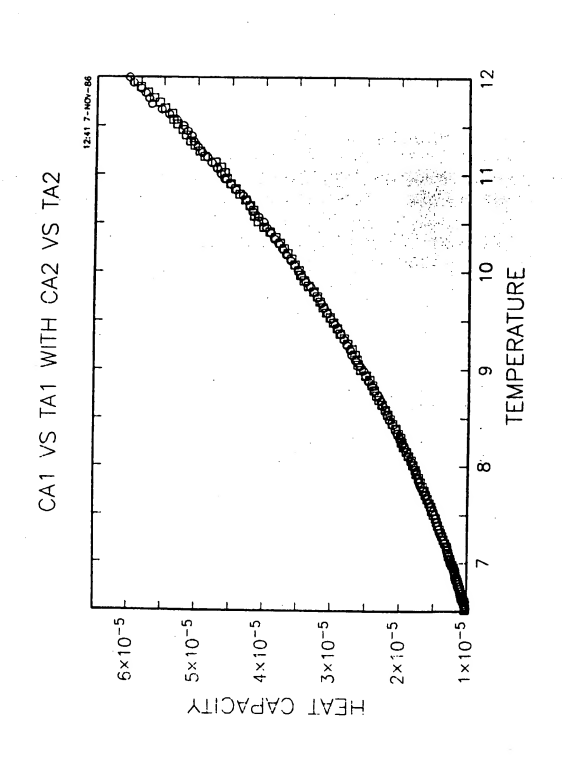


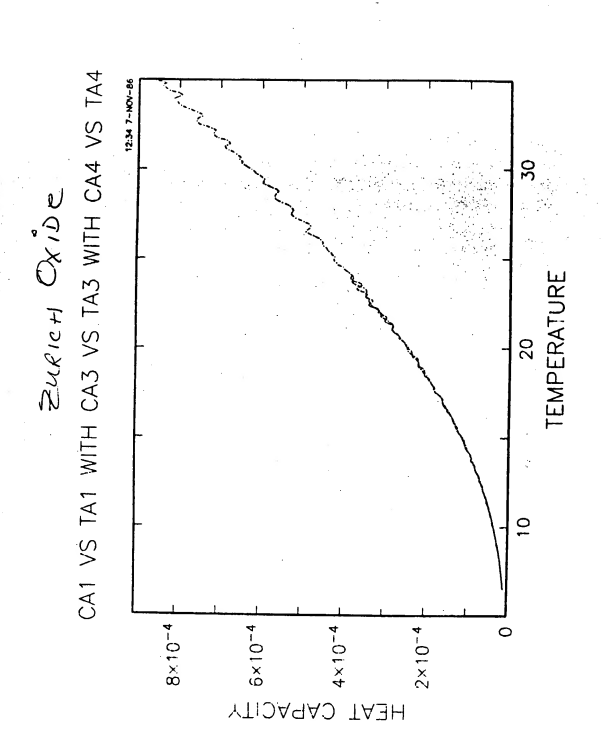


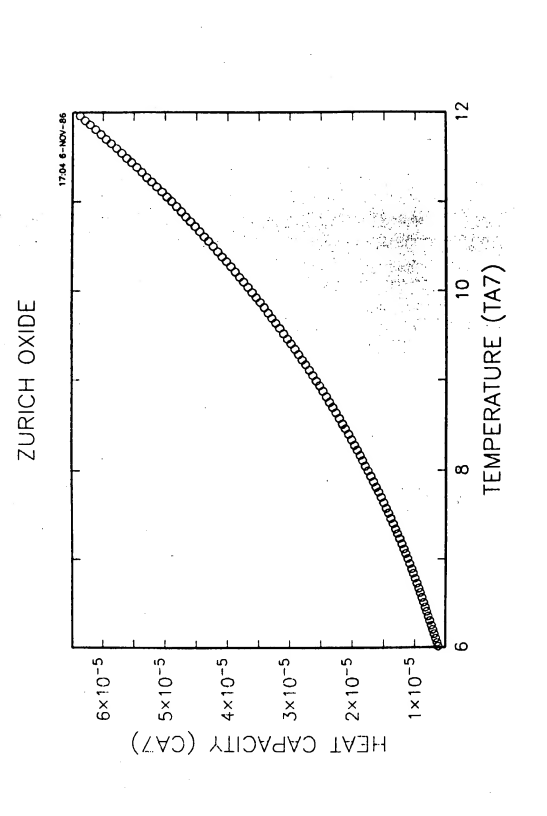


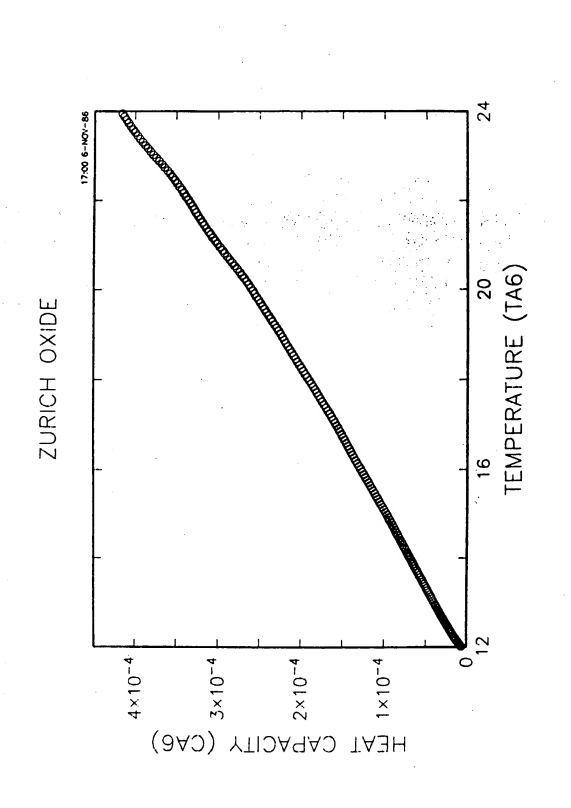


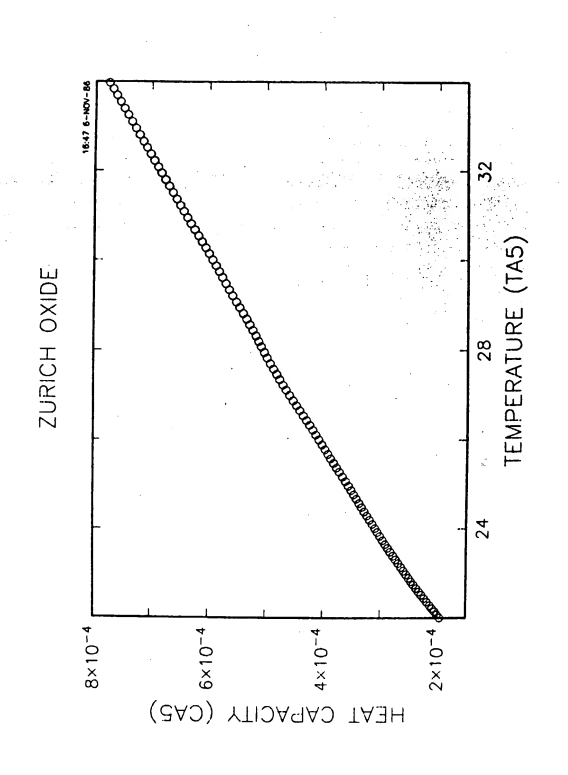






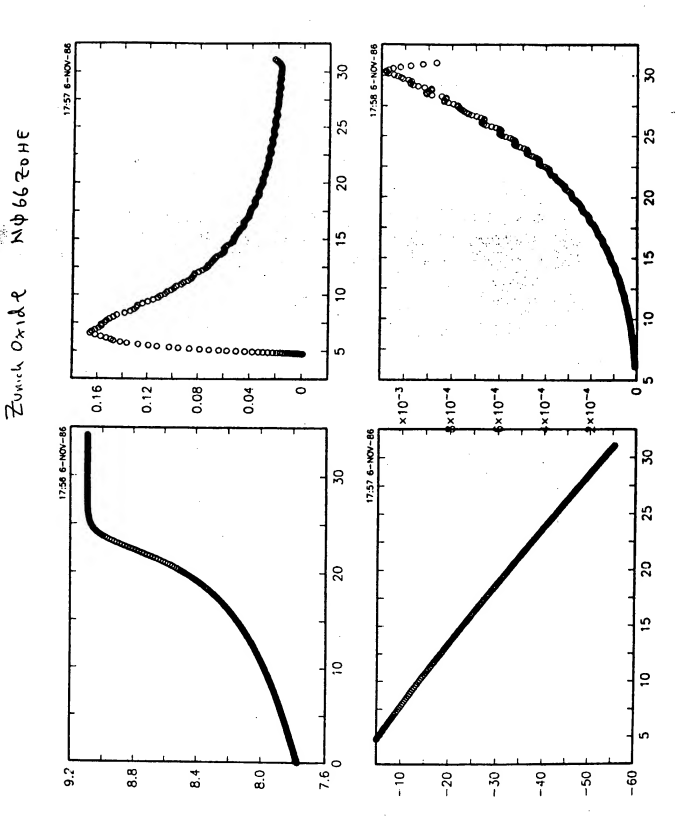


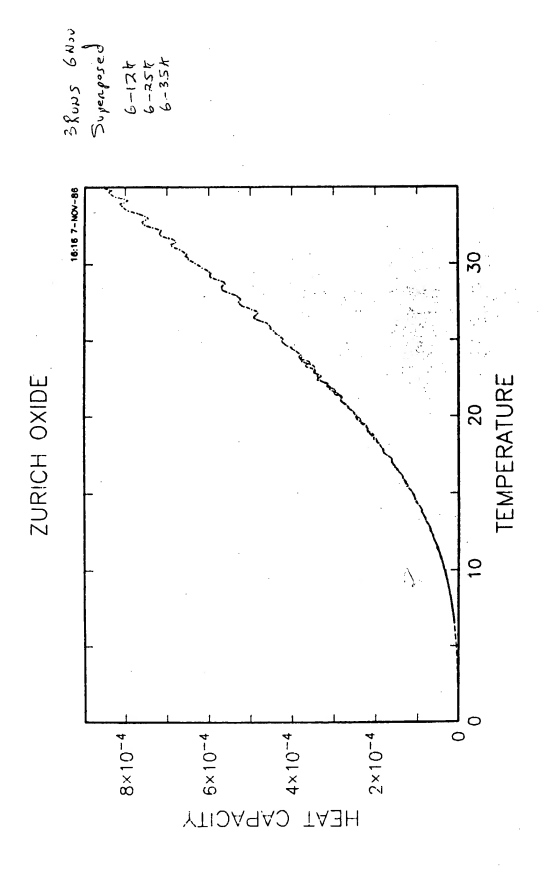




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